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ORIGINAL RESEARCH

Keeping an eye on pain: investigating visual attention biases in individuals with chronic pain using eye-tracking methodology

Samantha R Fashler Joel Katz

Department of Psychology, York University, Toronto, Ontario, Canada Abstract: Attentional biases to painful stimuli are evident in individuals with chronic pain, although the directional tendency of these biases (ie, toward or away from threat-related stimuli) remains unclear. This study used eye-tracking technology, a measure of visual attention, to evaluate the attentional patterns of individuals with and without chronic pain during exposure to injury-related and neutral pictures. Individuals with (N=51) and without chronic pain (N=62) completed a dot-probe task using injury-related and neutral pictures while their eye movements were recorded. Mixed-design analysis of variance evaluated the interaction between group (chronic pain, pain-free) and picture type (injury-related, neutral). Reaction time results showed that regardless of chronic pain status, participants responded faster to trials with neutral stimuli in comparison to trials that included injury-related pictures. Eye-tracking measures showed within-group differences whereby injury-related pictures received more frequent fixations and visits, as well as longer average visit durations. Between-group differences showed that individuals with chronic pain had fewer fixations and shorter average visit durations for all stimuli. An examination of how biases change over the time-course of stimulus presentation showed that during the late phase of attention, individuals with chronic pain had longer average gaze durations on injury pictures relative to pain-free individuals. The results show the advantage of incorporating eye-tracking methodology when examining attentional biases, and suggest future avenues of research.

Keywords: attentional biases, chronic pain, avoidance, hypervigilance, dot probe

The way individuals with pain attend to pain-related stimuli has been implicated in the development and maintenance of chronic pain.^{1–3} For example, the Schema Enmeshment Model of Pain¹ proposes that pain-, illness-, and self-schemas are overprocessed and that pain increases the personal relevance of pain and illness to the self, becoming enmeshed over time. The enmeshment of schemas may become problematic when a disproportionate amount of attention is directed toward pain-related stimuli,¹ implicating hypervigilance as the primary mechanism of maladaptive processing of pain-related information. However, evidence for attentional biases toward pain-related information in individuals with chronic pain is inconsistent:^{1,2,4–6} some investigations have found a bias toward threatening stimuli,^{7–9} away from threatening stimuli (later in attentional processing),^{10,11} or no evidence of a bias.^{4,12} These equivocal findings may be due to limitations in the measurement of attentional biases, stimulus differences (eg, words, pictures), or individual differences (eg, some individuals may exhibit avoidance or attending biases).

Correspondence: Joel Katz; Samantha R Fashler Department of Psychology, York University, 4700 Keele Street, BSB 232, Toronto, ON M3J IP3, Canada Tel +1 416 736 2100 Ex 40557 Fax +1 416 736 5814 Email jkatz@yorku.ca; sfashler@yorku.ca



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Previous investigations of attentional biases have been hampered by the limitations of the dot-probe task, a popular method for assessing selective attention.¹³ The dot-probe task uses reaction time (RT) as the primary outcome measure to detect attentional biases, which assumes that gaze location directly corresponds to motor behavior. Currently, more direct measures of visual attention are available, such as eye-tracking technologies, which can track real-time eye movements during attentional tasks. In this way, tracking eye movements offers a complementary measure of attention that may provide a better contextual understanding of visual biases. The type of stimuli (eg, words versus pictures) used to assess attentional bias may also contribute to the variability in study outcomes and the equivocal findings noted earlier.

Attention is a continuous process that changes over time. Visual attention phases represent periods of attentional processing that are qualitatively and temporally distinct. Although there is no consensus regarding what the attentional phases are (ie, their specific qualities or duration^{14,15}), they have been hypothesized to include the following in relation to a stimulus: orientation, engagement, maintenance, disengagement, re-engagement, and avoidance.^{2,3,16-20} It is possible that certain phases are important in distinct ways in the development and maintenance of attentional biases in individuals with chronic pain.² It is difficult to separately examine phases as there is likely a substantial overlap, with most research associating a specific time period with a specific attentional phase. As eye-tracking provides a continuous measure of visual attention, it permits the capture of attention at different visual attention phases over the course of a stimulus's presentation.

The present study used a dot-probe task coupled with eye-tracking technology to investigate attentional differences between individuals with and without chronic pain. We hypothesized that individuals with chronic pain would 1) respond faster on congruent dot-probe trials; 2) look more frequently at injury pictures; 3) display greater attentional maintenance to injury pictures; and 4) look for longer at injury pictures in later phases of attention than pain-free individuals.

Methods

The present study is part of a larger project investigating attentional biases in individuals reporting pain. The methods and data preparation are the same as those used by Fashler and Katz²¹ with the exception of the section on visual stimuli. The study protocol was approved by the Human Participants

Review Subcommittee at York University (ethics certificate number: 2013-048). Written informed consent was obtained from participants before beginning the study.

Participants

Recruitment

Participants were recruited through York University's Undergraduate Research Participant Pool. Before participants enroll in studies, they complete a prescreening survey composed of questions submitted by various faculty research laboratories. Responses to questions determine the studies that are visible to the student. For the present study, the question "do you experience physical pain on a regular basis?" was included in the prescreening survey. Of the 3,163 students who completed the prescreening survey, 655 (20.71%) positively endorsed the statement, 2,508 (79.29%) did not; 26 declined to respond. In order to recruit equal sample sizes of participants with and without pain, two identical study postings were created: one was visible to students who positively endorsed the prescreen question related to pain and the other was visible only to students who did not. This facilitated recruitment of equal sample sizes of participants with and without pain. Participants received one course credit for completing the study.

Demographic information

The final sample included 113 participants. The age range of the sample was 18–44 years, ($M_{age} = 21.32$, $SD_{age} = 4.35$). Eighty-four participants were female (74.30%) and 29 were male (25.70%). The sample was ethnically diverse, with participants self-identifying as Caucasian (32.54%), South Asian (28.57%), African (10.32%), East Asian (9.53%), Middle Eastern (8.73%), Hispanic/Latino (4.76%), Caribbean (3.97%), aboriginal (0.79%), and undisclosed (0.79%). Forty-nine (39.84%) participants wore vision-correction aids during testing, with 16 (14.20%) using contact lenses and 33 (29.20%) using glasses.

For the purpose of the present study, chronic pain was defined as the presence of ongoing pain that had persisted for 3 months or longer.²² According to this criterion, 51 (45.13%) participants reported chronic pain. Of these, seven (13.70%) participants reported experiencing pain for 3–6 months, five (9.80%) for 6–12 months, and 39 (76.50%) for 12 months or longer. Participants reported pain in one to five body locations ($M_{location} = 2.16$, SD_{location} =1.08): 30 reported neck and/or back pain, 22 reported headache/migraine pain, 21 reported ankle and/or knee pain, 15 reported shoulder pain, 12 reported stomach pain, five

reported hip pain, two reported arm pain, one reported eye pain, one reported jaw pain, and one declined to respond to this question. Most participants reported that the pain commenced after an injury (47.1%) or that they were unsure how the pain started (35.3%), while 7.8% of participants stated it started due to overuse and/or stress, 5.9% due to an illness, and one (0.79%) due to a hereditary predisposition. Frequency of pain was reported as daily (64.7%), weekly (33.3%), or other (2.0%) with the average intensity as mild (15.7%), moderate (74.5%), or severe (9.8%). When completing the questionnaires, 30 participants reported their present pain intensity as mild (73.3%) or moderate (26.6%) pain. Approximately half (52.9%) of chronic pain participants reported that they used pain medication on a regular basis. Of these participants, 63.0% used nonsteroidal anti-inflammatory drugs, 59.3% used acetaminophen, 7.4% used opioids, and 29.6% reported using other medications to manage their pain.

Participants reporting pain (N=51) did not differ significantly from those not reporting pain (N=62) on age, t (111) =0.71, P=0.705, sex, χ^2 (2, N=113) =0.001, P=0.969, use of contact lenses, χ^2 (1, N=113) =0.014, P=0.905, or glasses, χ^2 (1, N=113) =0.77, P=0.381. Pearson χ^2 2×4 analysis using pain status (chronic pain, pain-free) and ethnicity (African decent, Asian descent, Caucasian, other) identified a significant difference according to ethnicity, χ^2 (3, N=113) =18.93, P<0.001. Post hoc comparisons showed a significantly greater proportion of participants self-identifying as Caucasian among those reporting pain.

Materials

Hardware

Eye movements were tracked using a Tobii T60 XL eyetracker with a 24-inch widescreen monitor and a 60 Hz data rate (Tobii[®] Technology, Falls Church, VA, USA). The eye-tracker is integrated into the monitor, allowing participants to move their heads freely throughout the study. The Tobii T60 XL has an accuracy rate of 0.5° and $<0.03^{\circ}$ drift, reducing differences between what the participant is looking at and what the equipment is recording. Two Dell Precision T3400 Intel[®] CoreTM 2 Quad CPUs with 4 GB of RAM were interfaced to facilitate data collection. The display resolution for the dot-probe protocol was 800×600 pixels to optimize video capture and data transfer. The study took place in a windowless room in order to standardize the lighting for all participants.

Software

E-Prime 2.0 Professional (Psychology Software Tools, Inc., Sharpsburg, PA, USA) was used to design, display, collect, and prepare data for the dot-probe task.²³ E-Prime provides millisecond precision in the capture of RTs. The protocol was designed in E-Studio, data were combined with E-Merge, and data were prepared for export with E-DataAid. Tobii Studio Professional 2.0 captured participant eye movements with programmed time-sensitive markers (Tobii[®] Technology, 2010). E-Prime and Tobii Studio software programs were interfaced with E-Prime 2.0 extensions for Tobii (Psychology Software Tools, 2011). Data were analyzed with SPSS 20 (version 20.0, IBM Corporation, Armonk, NY, USA).

Visual stimuli

Sixty pictures were selected from the International Affective Picture System (IAPS)²⁴ to use as visual stimuli for the dotprobe task. IAPS is a collection of over 1,000 pictures that have normative affective ratings on two dimensions of emotion: valence (negative/aversive to positive/attractive) and arousal (calm to excited). Each trial in the dot-probe task consists of two pictures presented simultaneously. The picture pair consisted of either 1) an injury picture and a neutral picture (injury–neutral pair) or 2) two neutral control pictures (control pair). The final set of pictures consisted of 15 injury–neutral pairs and 15 neutral–neutral control pairs for a total of 60 pictures (30 pairs). Following selection of stimuli, the picture size of all pictures was standardized to 341×256 pixels.

Injury-neutral pairs

Fifteen injury-related IAPS pictures were selected (3030, 3103, 3180, 3181, 3185, 3195, 3211, 3213, 3215, 3220, 9042, 9405, 9415, 9592, and 9599). Inclusion criteria were 1) depiction of physical injury to a human (eg, needle insertion, black eye, open wound, burned skin, stitches); 2) low score for valence (ie, negative); and 3) high in arousal (ie, emotionally charged). Mean (SD) scores for arousal and valence were 5.76 (0.57) and 2.63 (0.65), respectively, for the injury pictures.

Fifteen neutral IAPS pictures were paired to the 15 injury IAPS pictures by depicting the same body region as the injury-related picture but without injury (eg, a picture displaying an injured hand was paired with a picture of a hand without any signs of injury) (2200, 2002, 2102, 2372, 2018, 2019, 2104, 7513, 7509, 2570, 2445, 2579, 2393, 2411, and 7092). Inclusion criteria were 1) no evidence of physical injury; 2) neutral valence (ie, neither positive nor negative);

and 3) low on arousal (ie, calm). Mean (SD) scores for arousal and valence were 3.57 (0.64) and 5.17 (0.55), respectively, for the neutral pictures. As expected, *t*-tests confirmed that in comparison to neutral pictures, injury pictures were significantly lower on valence (P<0.001) and significantly higher on arousal (P<0.001).

Control pairs

A set of 30 different control IAPS pictures was selected (7000, 7001, 7002, 7003, 7004, 7006, 7009, 7035, 7019, 7020, 7021, 7010, 7012, 7017, 7041, 5390, 7032, 7030, 7061, 7081, 7025, 7175, 7211, 7150, 7057, 7950, 7026, 7090, 7235, and 7052). Inclusion criteria were 1) an inanimate object (eg, household items, such as spoon and clock); 2) neutral valence (ie, neither positive nor negative); and 3) low on arousal (ie, calm). Mean (SD) scores for arousal and valence were 2.89 (0.62) and 5.05 (0.24), respectively, for all 30 control pictures. As expected, *t*-tests confirmed that the 15 pairs of control pictures did not differ significantly on valence or arousal (both P>0.450).

Behavioral measures

Reaction time measures

RT (in milliseconds) was recorded to capture the speed of participant responses to the location of the dot probe. For

each participant, a congruency index, incongruency index, and neutral comparison index were computed.

Congruent, incongruent, and neutral comparison indexes

These indexes are based on calculations by Koster et al²⁵ and Roelofs et al.²⁶ A trial is considered to be congruent when the dot probe appears in the location on the screen where the target picture (ie, injury picture) was presented and incongruent when the dot probe appears in the location of the neutral picture (Figure 1). The indexes are based on four mean RT scores, \overline{RT}_{tldr} , \overline{RT}_{trdr} , \overline{RT}_{trdl} , and \overline{RT}_{tldl} , which are computed by taking the mean of the 15 trials in each of the four conditions (tldr, trdr, trdl, and tldl) where "t" is the target (injury picture), "d" is the dot probe, "l" is the left position on the screen, and "r" is the right position on the screen.

The congruency index is calculated by taking the grand mean of the mean RTs of congruent trials that are presented on the right and left sides of the computer screen: $\overline{\text{RT}}_{\text{congruent}} = (\overline{\text{RT}}_{\text{tldl}} + \overline{\text{RT}}_{\text{trdr}})/2$. Similarly, the incongruent index is calculated by taking the grand mean of the mean RTs of the responses of incongruent trials that are presented on the right and left sides of the computer screen: $\overline{\text{RT}}_{\text{incongruent}} = (\overline{\text{RT}}_{\text{trdl}} + \overline{\text{RT}}_{\text{trdl}})/2$. The neutral comparison

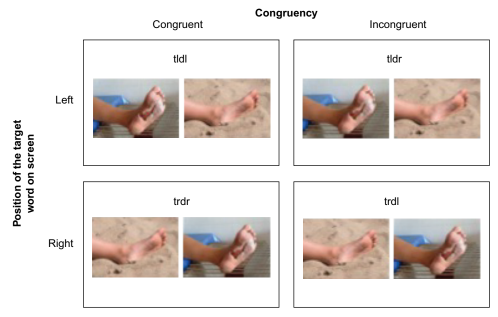


Figure I Visual depiction of counterbalanced trials of pictures presented during the dot-probe task.

Notes: Injury-related picture (target, t) on left (I) side of screen followed by the dot (d) on the left side of the screen (left, congruent or "tldl"); injury-related picture on left side of screen followed by the dot on the right (r) side of the screen (left, incongruent or "tldr"); injury-related picture on right side of screen followed by the dot on the right of the screen (right, congruent or "trdr"); and injury-related picture on right side of screen followed by the dot on the left side of the screen (right, congruent or "trdr"); and injury-related picture on right side of screen followed by the dot on the left side of the screen (right, incongruent or "trdr"). The images depicted are not drawn from the IAPS database but are similar to the stimuli used in the study protocol (photo credits: Mariusz Jurgielewicz © 123RF. com, Sarit Wuttisan © 123RF.com).

Abbreviation: IAPS, International Affective Picture System.

index is calculated to permit comparison with the congruent and incongruent indexes. It is calculated by taking the grand mean of the mean RTs for the neutral control picture pairs: $\overline{RT}_{neutral} = (\overline{RT}_{tldr} + \overline{RT}_{trdr} + \overline{RT}_{trdl} + \overline{RT}_{tldl})/4.$

Visual attention measures

Tobii I-VT fixation filter settings were used to classify fixations as they provide validated and robust function parameter values that are easily replicated for future research. As such, maximum gap interpolation (ie, merging) of fixations was set to 75 ms, the max angle between fixations was 0.5° , and fixations had to be >60 ms.²⁷ These settings provide accurate fixation classifications for most research.^{26,27} Additionally, data were screened for validity and only used if the data collected were associated with the correct eye.

Gaze patterns were recorded for the duration of the dotprobe task. Data were based on eye movements recorded within a region on the left and right sides of the screen defined by the experimenter as an "area of interest" (AOI). An AOI, 250 pixels in length by 107 pixels in height, was demarcated at mirror image regions on the center and midline of the left and right sides of the screen within which picture stimuli (injury-related, neutral, or control) were presented. Seven visual attention variables were calculated. Two measured the frequency of eye movements within an AOI: number of fixations assessed the number of times the participant's eyes focused on an injury or neutral picture and number of visits assessed the total number of times the gaze left an AOI (ie, a "visit" is defined by one or more contiguous eye movements within an AOI and ends when the eyes move outside the AOI). Two variables measured attentional maintenance within an AOI: mean fixation duration assessed the mean time associated with all fixations and mean visit duration assessed the mean time spent looking during all visits. Three variables measured the mean gaze duration per injury and neutral pictures according to three time intervals that spanned the 2,000 ms that the pictures appeared on the screen: early phase mean gaze duration (between 0 and 500 ms); middle phase mean gaze duration (between 500 and 1,000 ms); and late-phase mean gaze duration (between 1,000 and 2,000 ms).

Self-report measures

Demographics and Pain History Questionnaire

Participants were asked basic demographic questions and questions related to their pain history and current pain

complaints, including the duration, location, frequency, and cause of any pain.

Questionnaires

Participants completed nine questionnaires assessing health, anxiety, and pain-related variables. The Pain Disability Index²⁸ is a seven-item scale that assesses the degree of daily interference due to pain. The Pain Catastrophizing Scale²⁹ is a 13-item scale that assesses an individual's tendency to ruminate and magnify painful experiences. The Fear of Pain Questionnaire-III³⁰ is a 30-item scale that measures fear related to the experience of severe pain, minor pain, and medical pain. The Pain Vigilance and Awareness Questionnaire³¹ is a 16-item scale that assesses an individual's daily preoccupation with pain. The State-Trait Anxiety Inventory³² is a 20-item scale that assesses general levels of trait anxiety. The Anxiety Sensitivity Index-3³³ is a 16-item scale that assesses fear of anxiety-related sensations. The Health Anxiety Questionnaire³⁴ is a 21-item scale that measures daily functioning interference related to worry and preoccupation regarding health, illness, and death. The Illness Sensitivity Index-Short Version³⁵ is a six-item scale that assesses the fear of experiencing illness. The Sensitivity to Pain Traumatization Scale³⁶ is a 20-item scale that assesses anxiety-related cognitive, emotional, and behavioral reactions to pain that resemble symptoms of posttraumatic stress disorder. For all scales, higher scores are related to feelings of higher disability, anxiety, hypervigilance, or impairment.

Procedure

After arriving at the lab and providing informed consent, participants completed a computer-administered survey consisting of the ten questionnaires described earlier. The demographic and pain history questions appeared at the beginning of the survey for all participants. The order of the remaining questionnaires was randomized within participants using an online survey manager (Sona Systems, Bethesda, MD, USA). Participants were led to a windowless room to complete the dot-probe task. They were positioned ~60 cm in front of the screen with eyes leveled with the center of the screen. Five-point eye calibration was performed with Tobii Studio software to customize the eye-tracking specifications to each participant. The computer input was adjusted to run E-Prime from a second computer whereupon a second eye calibration was performed. Next, the dot-probe task was introduced and described. Participants were told that they were to fix their gaze at the central fixation cross, that a pair of pictures would appear briefly on the screen, and that a dot would be presented in the location of one of the pictures. Their task was to identify the location of the dot as quickly and accurately as possible. Participants then completed 12 practice trials to familiarize themselves with the task.

The protocol for each trial consisted of three parts: 1) a fixation cross ("+") appeared at the center of the screen for 500 ms; 2) two pictures (injury-neutral or neutral-neutral) were presented simultaneously, one on the left, and the other on the right side of the screen for 2,000 ms; and 3) immediately after the offset of the pictures, a dot probe (" \bullet ") appeared on the left or right side of the screen (in the prior location of the picture) for 2,000 ms or until the participant responded by pressing one of two keyboard keys. Participants were told to press the "P" with their right index finger if the dot probe was on the right side of the screen and the "Q" with their left index finger if the dot probe was on the left. All text and symbols appeared in black on a white background. Presentation of picture pairs was counterbalanced, controlling for picture type location (left, right) and dot-probe location (left, right), so that each picture pair was presented four times (Figure 1). In total, participants completed 120 trials (four trials each of the 15 injury-neutral picture pairs and four trials each of the 15 neutral-neutral picture pairs). The presentation order of trials was randomly generated by E-Prime (ie, the order selection was set to "random") for each participant.

Statistical analysis

Comparisons of nominal variables were made with X^2 tests of independence. Reaction time (RT) indexes were analyzed using a mixed-design 2×3 analysis of variance using group (chronic pain, pain-free) as the independent samples factor and index type (congruent, incongruent, neutral) as the within-subjects factor. Number of fixations and number of visits were each analyzed with a 2×2 mixed-design analysis of variances using group (chronic pain, pain-free) as the between-subjects factor and picture type (injury–neutral) as the within-subjects factor. Significant interactions were broken down with simple main effects analyses.

Results

Data preparation

Reaction time measures

Incorrect trials or trials that had RIs >2,000 ms were excluded from the calculation of the mean RI scores.²⁵ According to these criteria, 1.02% of trials were excluded from the calculation of the mean RI scores. Missing RIs scores were prorated if 80% or more of trials were available.^{37,38} One participant with insufficient data was excluded from the analysis of the RI data.

Visual attention measures

Eye-tracking recordings were screened for the quality of visual gaze capture. Of the 135 participants who were recruited, only those whose gaze capture exceeded 75% were included, consistent with previous eye-tracking research.³⁹ Lower gaze capture may be related to compromised attention, consistently lowered/closed eyelids, or reflections off glasses (eye-wear) that obscure the visual recording.⁴⁰ This led to the exclusion of 20 participants with poor capture ($M_{capture} = 60.90\%$, SD_{capture} =12.52%) and two participants who were unable to be calibrated. The mean eye capture rate for included participants (N=113) was 88.44%, SD =5.34%.

Participants included in the final analysis did not differ significantly from those with insufficient gaze data (N=22) on age, use of contact lenses, glasses, or presence of chronic pain (P>0.064 for all). A greater proportion of males (100%) than females (80%) had sufficient gaze data, χ^2 (1, N=134) =6.88, P=0.009. The omnibus X^2 analysis showed a significant effect according to ethnicity (African, Asian, Caucasian, other), c² (3, N=135) =11.690, P=0.009. Analysis of residuals showed that a significantly greater proportion of Asian and Caucasian participants had sufficient gaze data compared with African and other ethnicity participants (P<0.05).

Questionnaires

Missing questionnaire items were prorated if 80% or more of questions were available.^{37,38} One participant responded to fewer than 80% of the questions on the Anxiety Sensitivity Index, State-Trait Anxiety Inventory (state version), and Sensitivity to Pain Traumatization Scale, and was consequently excluded from the analysis of these questionnaires.

Reaction time measures

Mauchly's test of sphericity was significant, X^2 (2) =11.308, *P*=0.004. As the estimate of sphericity was >0.75, the Huynh–Feldt correction was used (Huynh and Feldt, 1976).⁴¹ Only the main effect of index type was significant, *F* (1.87, 207.29) =9.296, *P*<0.001, η_p^2 =0.077 (Table 1). Followup analyses showed that the neutral index (M =568.87,

Table I Reaction time index scores (in ms) for individuals with(N=51) and without chronic pain (N=62)

Index type	Chronic pain group Mean (SD)	Pain-free group Mean (SD)	
Congruency index (ms)	577.30 (126.81)	603.52 (127.86)	
Incongruency index (ms)	578.91 (116.17)	604.30 (141.89)	
Neutral comparison	560.12 (108.91)	577.63 (111.41)	
index (ms)			

Abbreviation: SD, standard deviation.

Visual attention measures	Chronic pain group	1	Pain-free group	
	Injury pictures Mean (SD)	Neutral pictures Mean (SD)	Injury pictures Mean (SD)	Neutral picture Mean (SD)
Frequency of gaze (N)				
Number of fixations	222.63 (56.49)	174.27 (37.51)	248.40 (61.78)	179.26 (41.45)
Number of visits	129.53 (24.18)	111.94 (20.90)	131.66 (25.69)	106.79 (18.84)
Attentional maintenance (S)				
Mean fixation duration	0.185 (0.03)	0.185 (0.04)	0.178 (0.02)	0.186 (0.03)
Mean visit duration	0.348 (0.06)	0.315 (0.06)	0.366 (0.06)	0.348 (0.06)
Attentional phases				
Early phase (0–500 ms)	0.136 (0.02)	0.142 (0.02)	0.132 (0.02)	0.138 (0.02)
Middle phase (500–1,000 ms)	0.155 (0.03)	0.153 (0.03)	0.158 (0.02)	0.149 (0.03)
Late phase (1,000–2,000 ms)	0.238 (0.07)	0.233 (0.07)	0.22 (0.04)	0.23 (0.05)

Abbreviations: SD, standard deviation; N, number; S, second

SD =10.43) was significantly shorter than both the congruency index (M =590.41, SD =12.04, P<0.001) and the incongruency index (M =591.60, SD =12.38, P=0.003).

Frequency of eye movements

For number of fixations, the main effect of picture type was significant, *F* (1, 111) =83.94, *P*<0.001, η_p^2 =0.431, with more fixations on injury pictures (M=235.52, standard error [SE] =5.62) than neutral pictures (M =176.66, SE =3.75). The main effect of group was also significant, *F* (1, 111) =4.79, *P*=0.031, η_p^2 =0.041, with fewer fixations overall by individuals with chronic pain (M =198.34, SE =5.24) than pain-free individuals (M =213.83, SE =4.76). For number of visits, the main effect of picture type was significant, *F* (1, 111) =76.58, *P*<0.001, η_p^2 =0.408, with more visits to injury pictures (M =130.60, SE =2.37) than neutral pictures (M =109.37, SE =1.87).

Maintenance of visual attention

For mean visit duration, the main effect of picture type, F(1, 111) = 15.26, P < 0.001, $\eta_p^2 = 0.121$, was significant with longer visits for injury pictures (M = 0.357, SE = 0.005) than neutral pictures (M = 0.325, SE = 0.007). The main effect of group was significant, F(1, 111) = 4.23, P = 0.042, $\eta_p^2 = 0.037$, with shorter visit durations for individuals with chronic pain (M = 0.332, SE = 0.007) than by pain-free individuals (M = 0.351, SE = 0.006). None of the effects were significant for mean fixation duration (Table 2).

Attentional phases

For early phase (0–500 ms) mean gaze duration, the main effect of picture type was significant, *F* (1, 111) =16.58, *P*<0.001, η_p^2 =0.130, with longer mean gaze duration on neutral pictures (M =0.140, SE =0.002) than injury pictures

Table 3 Comparison of questionnaire data for individuals with chronic pain (N=51) and without chronic pain (N=62)

Scale	Chronic	Pain-free	t (df)	P-value
	pain group	group		
	Mean (SD)	Mean (SD)		
PDI	20.18 (12.00)	5.37 (9.21)	–7.42 (92.4⊺)ª	<0.001*
PCS	19.22 (11.69)	16.16 (12.00)	-1.36 (111)	0.176
PVAQ	46.18 (10.69)	41.23 (9.64)	-2.59 (111)	0.011*
PASS-SF	37.72 (17.61)	36.03 (19.94)	-0.474 ()	0.637
CPAQ	77.57 (14.74)	66.57 (14.74)	-4.15 (111)	<0.001*
STAI-S	39.53 (12.39)	34.01 (11.14)	-2.49 (110)	0.014*
ASI-3	21.49 (12.86)	22.36 (13.33)	0.349 (110)	0.728
ISI-SF	16.71 (8.64)	16.23 (9.55)	-0.277 (111)	0.782
SPTS	27.91 (13.48)	26.93 (15.39)	-0.356 (110)	0.722

Notes: *Indicates a significant difference in scores at an adjusted P=0.01 to correct for multiple comparisons. ^aAdjusted t-test score reported due to a violation in Levene's test of equality of variance, F(2, 108) = 5.75, P=0.018.

Abbreviations: ASI-3, Anxiety Sensitivity Index-3; CPAQ, Chronic Pain Acceptance Questionnaire; df, degrees of freedom; ISI-SF, Illness Sensitivity Index-Short Version; PASS-SF, Pain Anxiety Symptoms Scale-Short Form; PCS, Pain Catastrophizing Scale; PDI, Pain Disability Index; PVAQ, Pain Vigilance and Awareness Questionnaire; SD, standard deviation; SPTS, Sensitivity to Pain Traumatization Scale; STAI-S, State-Trait Anxiety Inventory (state version).

(M =0.134, SE =0.002). For middle phase (500–1,000 ms) mean gaze duration, the main effect of picture type was also significant, F(1, 111) = 7.81, P=0.006, $\eta_p^2 = 0.066$, but in this case, there was a longer mean gaze duration on injury pictures (M =0.157, SE =0.002) than neutral pictures (M =0.151, SE =0.003). For late-phase mean gaze duration, the picture by group interaction was significant, F(1, 111) = 6.46, P=0.012, $\eta_p^2 = 0.055$. Simple main effects analysis showed that for injury pictures, individuals with chronic pain (M =0.238, SD =0.067) had significantly longer mean gaze durations than pain-free individuals (M =0.217, SD =0.035, P=0.038).

Group differences

Participants with and without pain were compared on questionnaire data (Table 3). Individuals with chronic pain reported

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significantly great pain disability (P<0.001), acceptance of chronic pain (P<0.001), state levels of anxiety (P=0.014), and pain vigilance (P=0.011). To examine the possible effect of pain medication use on task performance and eye movements, individuals with chronic pain who reported frequently using pain medications were compared with those who did not. No significant group or interaction differences were found according to RI, eye movements, or psychological variables.

Discussion

The purpose of this study was to investigate patterns of visual attention biases to threat-related pictures of injuries in individuals with and without chronic pain using two measures of biases, RI and eye-tracking methodology. RI measures showed that participants responded faster on neutral trials in comparison to congruent or incongruent trials regardless of chronic pain status. Likewise, eye-tracking measures showed that participants displayed an overall bias toward looking at injury-related pictures based on frequency of looks (including both fixations and visits) and attentional maintenance (including mean visit duration) but this did not differ among those with or without chronic pain. An examination of results within the three phases of attentional processing showed longer mean total gaze duration toward neutral and injury pictures during the early and middle phases, respectively, regardless of chronic pain status. In contrast, during the late phase of attention, individuals with chronic pain, but not the pain-free controls, displayed longer mean total gaze duration to injury-related than neutral pictures. The sample additionally showed unsurprising differences according to pain status for several psychological measures: pain disability, pain acceptance, state levels of anxiety, and pain vigilance were significantly higher for individuals with chronic pain. Together, the present results highlight the advantages of monitoring eye movements over simple RI to measure painrelated visual attention. Moreover, they provide novel data pointing to a pattern of late-phase attentional hypervigilance to injury-related pictures for individuals with chronic pain.

Until recently, the primary measure of attentional processing used in the dot-probe paradigm has been RI.¹³ Specifically, it predicts that faster RIs on congruent trials compared to neutral trials indicate vigilance to threat, which was not supported by the results of the present study, and that slower RIs on incongruent trials compared to neutral trials indicate difficulty disengaging from threat, which was supported in the present study.²⁵ Difficulty disengaging attention from a threat occurs when there is a struggle to redirect attention after a stimulus has been processed⁴² and this has been proposed to describe the nature of the attentional bias in anxiety,^{25,43,44} pain catastrophization,¹⁸ and rheumatoid arthritis.⁴⁵ It is not possible to determine with certainty, based on RI alone, whether participants in the present study had difficulty disengaging attention or were actively engaged. The delay on trials containing injury-related images may have been related to a negative emotional reaction, distraction (ie, full attention toward something else was impaired), or curiosity. Distinguishing between types of attentional biases, such as hypervigilance and difficulty disengaging from threat, is important from both a theoretical and clinical perspective. Knowing the nature of the bias will help improve our understanding of possible maladaptive attentional processing or dysfunction that, in turn, may inform effective treatment strategies. It is also noteworthy that we did not find a significant interaction effect for RI variables, but as described earlier, this was the case with eye-tracking variables. This shows that eye-tracking measures can be more sensitive than RI in identifying differences in attentional processes between individuals with and without chronic pain, demonstrating its utility as a complementary form of measurement of attentional biases.

Measures of eye-tracking frequency showed that injuryrelated pictures received more frequent fixations and visits than other neutral pictures. Although this provides support for the notion that humans are evolutionarily predisposed to attend to threatening stimuli in the environment,⁴⁶ it does not provide support for the Schema Enmeshment Model of Pain, which predicts that individuals with pain will show increased attentional processing of injury-related stimuli.1 In fact, there was evidence of fewer fixations for individuals with chronic pain. Although the absence of an interaction effect shows that attention was increased for both injury- and neutral-related pictures, this may indicate a pattern of overall avoidance. A similar pattern emerged for measures of attentional maintenance. Although no findings were significant for mean fixation duration, all participants had longer mean visit durations on injury-related pictures, with individuals with chronic pain looking at all pictures for shorter sustained period of time.

The use of eye-tracking in the present study permitted novel analysis that assessed how visual attention biases may differ across visual attention phases. The orientation phase of attention is considered to be the initial period of attentional activation toward a new stimulus and hypothesized to occur before the first 500 ms of looking at a stimulus.^{3,20,47} In the present study, during the orientation phase, the results showed that regardless of chronic pain status, participants had longer mean gaze durations to neutral pictures compared to injury pictures. This finding adds to the mixed evidence of previous RT-based dot-probe investigations regarding biases in early stages of processing,³ including toward threat,⁸ away from threat,¹⁰ or no effect.^{48,49} Interestingly, during the middle phase of attentional processing (500-1,000 ms), the total gaze duration shifted toward injury-related pictures, providing evidence that the nature of attentional patterns can change across attentional phases (ie, orientation versus maintenance) and supporting the frequency and maintenance findings earlier. During the late phase of attention (1,000–2,000 ms), the significant interaction showed that individuals with chronic pain had a longer mean gaze duration to injury-related pictures than did pain-free individuals. This differential pattern of attention over time supports the findings of a recent metaanalysis by Schoth et al³ of ten studies that used the visual dot probe. They found only a modest effect (Hedge's adjusted g=0.29) for orientation attentional biases in individuals with chronic pain (stimulus presentation time between 300 and 500 ms), which was more pronounced for maintained attention (stimulus presentation time of 1,250 ms; Hedges' adjusted g=0.42). Similar conclusions were drawn from a recent meta-analysis by Crombez et al² examining the effect of subliminal and supraliminal exposure durations, with the authors concluding that "conscious and elaborative processes are critical for attentional biases to emerge".

The present study has a number of limitations. Eyetracking methodology relies on the assumption that eye movements reflect visual attention. However, visual attention can occur in the absence of eye movements.^{50,51} This effect may be accentuated when exposed to a distracting task, leading to "inattentional blindness", whereby individuals may not recall exposure to a stimulus even when they looked directly at it.52,53 Therefore, although tracking eye movements is considered a strong indicator of visual attention,^{51,54} it is not a perfect reflection of overt attentional engagement. This study used a sample of undergraduate students who likely experience pain and disability that is not as severe as clinical samples.⁷ Therefore, further studies should evaluate attentional patterns in clinical populations. Finally, the present study was crosssectional, evaluating differences between individuals with and without pain rather than how specific attentional patterns may predict pain. Longitudinal study designs are needed to establish how attention may contribute to the development and maintenance of chronic pain.

This study examined attentional biases in individuals with and without chronic pain. Using eye-tracking technology provided more direct measures of visual attention than traditional dot-probe investigations. We found that overall,

individuals with and without pain responded faster to neutral stimuli contrary to the assumptions of the dot-probe paradigm, implying a difficulty to disengage from injuryrelated pictures. On measures of frequency and attentional maintenance, participants showed greater attention toward injury-related pictures regardless of pain status, and individuals with chronic pain had fewer fixations and shorter average visit durations than those without pain. The pattern of attentional biases changed over the course of stimuli presentation, showing initial avoidance of injury-related pictures followed by increased vigilance. During late-phase attention, only individuals with pain showed a residual bias toward injury-related pictures. Taken together, the novel use of examining attention over time with eye-tracking technology provides insight into how visual attention biases can change over time. Further research is needed to validate the present findings with clinical samples and with a longitudinal design.

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Disclosure

The authors report no conflicts of interest in this work.

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