Generalization of Pain-Related Fear Based on Conceptual Knowledge

Ann Meulders
Kristof Vandael
University of Leuven
Johan W.S. Vlaeyen
University of Leuven
Maastricht University

Increasing evidence suggests that pain-related fear is key to the transition from acute to chronic pain. Previous research has shown that perceptual similarity with a pain-associated movement fosters the generalization of fear to novel movements. Perceptual generalization of pain-related fear is adaptive as it enables individuals to extrapolate the threat value of one movement to another without the necessity to learn anew. However, excessive spreading of fear to safe movements may become maladaptive and may lead to sustained anxiety, dysfunctional avoidance behaviors, and severe disability. A hallmark of human cognition is the ability to extract conceptual knowledge from a learning episode as well. Although this conceptual pathway may be important to understand fear generalization in chronic pain, research on this topic is lacking. We investigated acquisition and generalization of concept-based pain-related fear. During acquisition, unique exemplars of one action category (CS+; e.g., opening boxes) were followed by pain, whereas exemplars of another action category (CS-; e.g., closing boxes) were not. Subsequently, spreading of pain-related fear to novel exemplars of both action categories was tested. Participants learned to expect the pain to occur and reported more pain-related fear to the exemplars of the CS+ category compared with those of the CS- category. During generalization, fear and expectancy generalized to novel exemplars of the CS+ category, but not to the CS- category. This pattern was not corroborated in the eyeblink startle measures. This is the first study that demonstrates that pain-related fear can be acquired and generalized based on conceptual knowledge.

Keywords: pain-related fear; acquisition; generalization; category-based learning; associative learning

The ability to learn which stimuli in the environment signal threat has an important adaptive advantage, because it enables us to initiate appropriate defensive responses that protect us from future harm (Vlaeyen, 2015). Nevertheless, adaptive learners face the challenge of how to deal with variations in the appearances of signaling stimuli. Stimulus generalization (Honig & Urcuioli, 1981; Kalish, 1969) allows individuals to extrapolate the predictive value of one stimulus to perceptually similar stimuli and minimizes the necessity to learn everything anew. As a consequence, the advanced capacity to detect similarities between unique but related stimuli fosters swift adaptation to a dynamic environment.

Pain is a vital motivator in learning because it typically alerts the individual of impending or actual bodily threat. It has been shown that associative learning is crucially involved in the acquisition of fear of movement-related pain (Meulders, Vansteenwegen, & Vlaeyen, 2011). In particular, using a voluntary joystick movement (VJM)
paradigm, Meulders and colleagues (2011) showed that after (repeated) pairing of a painful electrocutaneous stimulus (unconditioned stimulus [pain-US]) with an initially neutral joystick movement (conditioned stimulus [CS]), this movement (CS+) elicited protective responses such as fear and avoidance (conditioned response [CR]), whereas another neutral joystick movement (CS-) that was not paired with pain did not evoke such CRs. Differential pain-related fear learning was apparent in self-reports and psychophysiological measures (startle eyeblink measure), as well as behavior (movement-onset latency)—that is, participants reported more fear in response to the CS+ than to the CS-. However, when the CS+ was present, the diagonal movements to the left (GS+) elicited more fear measured with both self-report and startle amplitude than did the diagonal movements lying between the CS+ and the CS-. The results corroborate previous findings at least in the startle eyeblink measures. More specifically, we showed a pain-related fear generalization gradient—that is, there was a linear decrease in startle amplitudes for GSs approaching the original CS-. Another study using a left-right hand judgment conditioning paradigm with pictures of hand postures as CSs and a painful electrocutaneous stimulus as the pain-US showed a similar gradient at least in the verbal measures (fear and expectancy; Meulders, Harvie, Moseley, & Vlaeyen, 2015). In this study, one hand posture (CS+; e.g., extreme hand flexion) was consistently paired with pain and another hand posture (CS-; e.g., extreme hand extension) was not. During generalization, we tested the spreading of fear and US-expectancy to a set of novel hand postures (GSs) with six grades of perceptual similarity to the CS+. Results confirmed that novel GSs that were more similar to the CS+ triggered more pain-related fear and US-expectancy than did those more similar to the CS-. Taken together, accumulating evidence suggests that pain-related fear can spread based on perceptual similarity.

A certain degree of fear generalization is adaptive, but excessive generalization to technically safe stimuli may become maladaptive and pathological. In a replication of Meulders and Vlaeyen (2013), we demonstrated that patients with fibromyalgia syndrome do not show selective fear generalization like healthy pain-free controls, but overgeneralize their fear to all novel (diagonal) movements (GSs). In a hand pain scenario contingency learning task with hand postures as cues and the words “pain” and “no pain” as outcomes, participants were asked to rate the likelihood that a fictive hand pain patient would feel pain when moving the hand into certain postures (Meulders et al., 2014). One hand posture (e.g., extreme hand flexion) was consistently paired with the word “pain” and another hand posture (e.g., extreme hand extension) was paired with “no pain.” During generalization, we tested the spreading of pain expectancy to a set of novel hand postures (GSs) with six grades of perceptual similarity to the pain-associated hand posture. Unilateral chronic hand pain patients showed flatter, asymmetric generalization gradients compared with healthy pain-free controls, with higher pain expectancy ratings for novel postures that were more similar to the CS+. At the CS+ side of the gradient, pain expectancy ratings did not differ between patients and controls, indicating a lack of safety learning rather than excessive fear in response to actual threat.
Taken together, accumulating evidence suggests that adaptive pain-related fear generalization based on perceptual similarity occurs in a healthy population, but that maladaptive overgeneralization characterizes chronic pain populations.

Humans also possess the ability to abstract conceptual details of a learning episode, which allows them to generalize conditioned fear behavior between physically dissimilar stimuli that are semantically related, a process known as category-based fear conditioning (Dunsmoor & Murphy, 2015). The involvement of higher-order cognitions (i.e., cognitions requiring a certain degree of abstract thinking) such as conceptual knowledge or category membership in fear generalization has been demonstrated by Dunsmoor, Martin, and LaBar (2013). Typically, the term “exemplars” is used to refer to low-level, specific members (e.g., hammer) belonging to a superordinate category (e.g., tools) that signifies a higher, more abstract level within a classification system. In the Dunsmoor et al. (2012) study, 50% of the basic-level exemplars of one superordinate category (i.e., tools) were paired with an aversive shock, whereas exemplars of another category (i.e., animals) were not. The results showed category-specific anticipatory skin conductance responses and shock expectancy ratings. In another study, the investigators showed category-specific increases in brain activity in the visual cortex and fear-learning networks, further corroborating the idea that humans can learn to associate fear with an entire category despite the heterogeneity in the physical appearance of stimuli (Dunsmoor, Kragel, Martin, & LaBar, 2013). Based on the parallels between pain-related fear in chronic pain conditions and other types of pathological fear in anxiety disorders, concept-based fear learning might affect the spreading of fear and avoidance in pain conditions as well (Dymond, Dunsmoor, Vervliet, Roche, & Hermans, 2015). If the person in our previous example experienced a shooting pain in the back, but rather than associate it with a specific movement during the yoga class, attributed it to yoga as a whole (an activity that according to this person belongs to the superordinate category “sports”), he or she may become afraid of different types of sports (i.e., exemplars belonging to the category “sports”; e.g., swimming, cycling, running) regardless of their perceptual resemblance to yoga, and as a consequence might cease all sport-related activities.

Research on this conceptual path of fear generalization is scarce, and to our knowledge this question has never been addressed in the field of pain-related fear. Therefore, we sought to investigate (a) acquisition of fear of movement-related pain to physically dissimilar exemplars of one superordinate “action” category (CS+), and (b) fear generalization to novel CS+ exemplars (GSs) belonging to the same conceptual category. We adapted the VJM paradigm using unique exemplars of superordinate action categories (opening/closing boxes) as CSs, and a painful electrocutaneous stimulus as the pain-US. Another set of unique exemplars (boxes with different combinations of color, size, and shape) of both action categories served as GSs. Conditioned pain-related fear was measured through self-reports (pain-related fear and US-expectancy ratings) as well as a psychophysiological index of fear (eyeblink startle response). We hypothesized that (a) at the end of acquisition, fear and US-expectancy ratings as well as the eyeblink startle responses would be higher for the exemplars of the CS+ category than for the exemplars of the CS- category; and (b) during the test of generalization, fear and US-expectancy ratings as well as the eyeblink startle responses would be higher for novel exemplars of the CS+ category (GS+) than for novel exemplars of the CS- category (GS-).

Method

Participants

In total, 50 healthy, pain-free individuals (31 women, 19 men) voluntarily participated in this study and received course credit or €12 (approximately $13) to compensate them for their time and effort. They were between 18 and 38 years old (M = 23.32, SD = 4.32). Participants were recruited using the departmental experiment management system and through word of mouth. Exclusion criteria were pregnancy, cardiovascular diseases, chronic or acute respiratory diseases (e.g., asthma, bronchitis), neurological diseases (e.g., epilepsy), any other severe medical condition, any current or past psychiatric disorders including clinical depression and anxiety disorders, chronic pain, uncorrected hearing problems, painful hand-related problems, the presence of a cardiac pacemaker or any other electronic medical devices, and any condition that might influence the ability to make judgments/give verbal ratings (e.g., cognitive impairment due to stroke or brain injury). All participants completed a checklist to ensure that none of these criteria were applicable. Next, they signed the informed consent form. The experimental protocol was approved by the Social and Societal Ethics Committee of the University of Leuven (registration number G-2015 01 147).

Stimulus Material and Measures

The experiment was run on a Windows XP computer (Dell Optiplex 755) with 2 GB RAM and an Intel Core2 Duo processor at 2.33 GHz and an ATI Radeon 2400 graphics card with 256 MB of video RAM. Stimulus presentation was controlled with the
free software package Affect 4.0 (Spruyt, Clarysse, Vansteenwegen, Baeyens, & Hermans, 2010).

CSs consisted of 20 unique exemplars of two functional categories: closing and opening boxes (10 opening boxes, 10 closing boxes). These boxes had different colors, shapes, and sizes to reduce the influence of mere perceptual similarity of the exemplars belonging to the same category. At the beginning of a trial, these boxes were either open or closed. In order to “open” a closed box or to “close” an open box, participants moved a (Logitech Attack 3) joystick into the signaled direction (i.e., to the left or to the right). During this movement, an animation was run showing the actual opening or closing of the box. The GSs were 16 novel and unique exemplars of the two learned functional categories (8 opening boxes, 8 closing boxes). Thus, these boxes had different color, shape, and size combinations than those used during the acquisition phase. An electrocutaneous stimulus (2-ms duration) delivered by a commercial constant current stimulator (DS7A, Digitimer, Welwyn Garden City, UK) served as the unconditioned stimulus (pain-US). This stimulation was administered through surface Sensormedics electrodes (8 mm) filled with K-Y gel that were attached to the wrist of the dominant hand. During the calibration procedure, participants received a series of electrocutaneous stimuli of increasing intensity and were asked to indicate how intense/painful each stimulus was on a scale from 1 to 10 (1 = You feel something but this is not painful, it is merely a sensation; 2 = This sensation starts to be painful, but it is still a very weak pain; 10 = This is the worst pain you can imagine). Participants were instructed that a subjective stimulus intensity of 8 (Significantly painful and demanding some effort to tolerate) was targeted, but were asked to notify the experimenter when they did not want to receive a stimulus of higher intensity or when they wanted the intensity to be set back to a lower level. Lowering the intensity of the electrocutaneous stimulus was allowed only during the calibration phase; during the actual conditioning procedure, the intensity of the selected pain-US was never adjusted. The mean subjective stimulus intensity was 8.00 (SD = 0.40, range = 6–9), and the mean physical stimulus intensity was 32.12 mA (SD = 20.47, range = 6–100).

Conditioned pain-related fear was measured through self-reports as well as a psychophysiological measure of fear learning—that is, the eyeblink startle response. The eyeblink startle response is a component of the reflexive cross-species, full-body defensive response mobilization, which is triggered by startle-evoking stimuli (e.g., acoustic startle probe) and can be measured by the tension in the muscles underneath the eye (Blumenthal et al., 2005; Davis, Walker, Miles, & Grillon, 2010). Startle modulation refers to the potentiation of the startle reflex during fear states elicited by the anticipation of an aversive stimulus (e.g., an electrocutaneous stimulus); fear-potentiated startle (FPS) has been widely used as a proxy measure of conditioned fear (Grillon, 2002; Lang, Davis, & Ohman, 2000). In the present setup, the startle probe was a 100 dBA burst of white noise with instantaneous rise time presented binaurally for 50 ms through headphones (Philips SHP2500). Startle eyeblink responses elicited by startle probes delivered during the CS/GS movements served as an index of cued pain-related fear. Startle eyeblink responses elicited by startle probes during the intertrial interval (ITI) served as an index of contextual pain-related fear (Vansteenwegen, Iberico, Vervliet, Marescau, & Hermans, 2008). We expected that the ITI startle responses would not be potentiated, because during this period it was safe and no painful electrocutaneous stimulus was anticipated. Therefore, in the present design, the ITI startle responses served as a baseline/control measure of psychophysiological responding.

PROCEDURE
We adapted the VJM paradigm (Meulders et al., 2011; Meulders & Vlaeyen, 2013) to study the acquisition and generalization of pain-related fear based on conceptual knowledge. The experiment was conducted during a 90-minute session and consisted of a preparation phase, a practice phase, a startle habituation phase, an acquisition phase, and a generalization test. CSs were unique exemplars of two functional, superordinate categories (opening/closing boxes). During the acquisition phase, exemplars of the CS+ category were followed by the pain-US in 80% of the trials, and exemplars of the CS- category were never followed by the pain-US. The functional category that served as the CS+ and CS- was counterbalanced across participants. This means that half of the participants received a painful stimulus paired with opening boxes, whereas the other half received a painful stimulus paired with closing boxes. Because the acquisition and generalization (tested under extinction) of category-based pain-related fear was under investigation, we used a partial (80%) reinforcement schedule, as it is known to slow down both acquisition (Dunsamo, Bandettini, & Knight, 2007; Schurr & Runquist, 1973) and extinction rates (Haselgrove, Aydin, & Pearce, 2004; Schurr & Runquist, 1973). In half of the trials, participants moved the joystick to the left to “open” a closed box (e.g., CS+ category) or to “close” an open box (e.g., CS- category), and in the other half of the trials participants moved the joystick to the right. That way,
the same movements (i.e., moving the joystick to the left) could lead to different outcomes (opening or closing boxes) and thus served a different purpose. Therefore, participants should learn that it is not the movement direction, the color, the shape, or the size of the boxes that predicts the pain-US but that it is the function of the behavior (i.e., opening or closing boxes) that predicts the pain-US. During the test of generalization, novel exemplars of the same functional categories (GSs)—that is, novel boxes with unique combinations of shape, color, and size other than those used in the acquisition phase—were tested.

*Preparation*
Upon arrival to the laboratory, participants were informed that the experiment involved the repeated presentation of electrocutaneous stimuli (pain-US) and short loud noises (acoustic startle probes). Participants were also told that they were free to decline participation at any time without any negative consequences. After providing informed consent, electrodes for eyeblink startle responses and the administration of the electrocutaneous stimulus were attached and the calibration procedure of the pain-US was initiated.

*Practice*
Before starting the practice phase, participants received extensive written instructions about the experimental task. These instructions were presented via Affect 4.0 (Spruyt et al., 2010) on the computer screen in front of them. During the practice phase, participants were requested to “open” or “close” boxes by moving the joystick into the direction signaled by a red asterisk (i.e., the *direction signal*). They could start moving when prompted by a white fixation cross that was presented in the middle of the computer screen (i.e., the *starting signal*; see Figure 1). In total, 12 trials were run: participants were requested to open six boxes and close six boxes with three joystick movements to the left and three joystick movements to the right in each CS category (opening/closing boxes). The presentation order of the trials was semirandomized with no more than two consecutive trials of the same CS category into the same movement direction.

Each trial consisted of a pre-CS interval of 3 s and a post-CS interval of 5 s. After the pre-CS interval, an open or closed box was presented in the middle of the screen. Either on the left or right side of the box, the direction signal appeared, indicating the direction of the joystick movement that needed to be performed. After 500 ms, the direction signal disappeared and questions assessing pain-related fear and pain-US expectancy were presented on the screen. Two seconds after answering these questions, the starting signal was presented. Upon presentation of the starting signal, participants were requested to “open” or “close” the presented box by moving the joystick as fast and accurately as possible into the signaled direction. During the movement, an

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**FIGURE 1** Schematic overview of the experimental task.
animation of the box closing/opening was shown, and when the movement was successfully completed, the box disappeared from the screen and a post-CS interval of 5 s elapsed before a new trial commenced. When participants moved in the wrong direction, an error message was shown to inform them, and the trial was restarted. During the practice phase, no acoustic startle probes or pain-USs were presented, and the experimenter gave online verbal feedback about the task performance.

**Startle Habitation**
Because it is expected that the first responses to startle probes are significantly larger than latter ones, we inserted a startle probe habituation phase to correct for such a possible confound in the data. In other words, initial responses to startle probes are disproportionately high (due to orienting responses and novelty), and decrease quickly (i.e., habituate) following a couple of probe presentations. Because modulation of the response was under investigation, we used the habituated startle response as an onset reference for the measures. This phase consisted of eight trials, each lasting 13 s (with a variable ITI of on average 2 s). On each trial, one startle probe was presented randomly between 8 and 12 s after trial onset. During this phase, the lights were dimmed, participants wore headphones, and no pain-USs were delivered.

**Acquisition**
This phase was similar to the practice phase, except that pain-USs and startle probes were now presented. Pain-related fear conditioning occurred over four acquisition blocks that each included the same 10 unique exemplars of both CS categories (10 open and 10 closed boxes; 80 trials in total). Again, half of the open boxes were to be closed with joystick movements to the right and the other half with joystick movements to the left, whereas half of the closed boxes were to be opened with joystick movements to the right and the other half with joystick movements to the left. For each participant, one functional category (e.g., closing boxes) was designated the CS+, and 80% of the exemplars belonging to this category were randomly reinforced with delivery of the pain-US after completing the signaled joystick movement. There was no contingency between the occurrence of the pain-US and the direction of joystick movement. The other functional category (e.g., opening boxes) served as the CS-, and none of the exemplars belonging to this category were reinforced with a pain-US. In 40% of the trials (i.e., 4 out of 10 exemplars of each CS category in each acquisition block), questions assessing pain-US expectancy and how afraid participants were to perform the movement were presented before they actually carried out the joystick movement. Questions were presented pseudo-randomly with the restriction that left/right joystick movements were equally distributed across CS+ and CS- assessment trials. We used 40% of assessment trials in order to follow the development of the CR without interfering too much with the conditioning process itself. After each acquisition block, participants also rated the pain-US intensity and unpleasantness and how afraid they were of the “relevant” CS categories (opening/closing boxes) as well as other “irrelevant” perceptual and proprioceptive features: (a) movement direction (left/right), (b) size (small/big boxes), and (c) color (dark/light-colored boxes).

Although a CS movement was of variable length depending on participants’ movement speed, a trial again included an ITI consisting of a pre-CS interval of 3 s and a post-CS interval of 5 s. The rest of the timing was the same as during the practice phase. On each trial, one startle probe was presented; within each acquisition block, six exemplars of the CS+ category and six exemplars of the CS- category were probed (for both categories: three during movement to the left, three during movement to the right). These startle probes were presented 200 ms after participants started “opening” or “closing” the boxes (i.e., when they started to move the joystick). Eight probes were presented during the ITI (four on CS+ trials and four on CS- trials); two probes were presented randomly between 700 and 2200 ms of the pre-CS interval, and two probes were presented randomly between 2200 and 3700 ms of the post-CS interval. Thus, in each acquisition block, 12 probes were presented during movements and 8 were presented during the ITI (i.e., context alone). Participants were not informed about the contingencies between the functional categories (CSs) and the pain-US.

**Test of Generalization**
The procedure for the generalization phase was similar to the acquisition phase. In this phase, we tested 8 novel exemplars of both the CS+ and CS- categories (respectively referred to as GS+ and GS-; 16 trials in total). These GSs were never followed by the pain-US. To prevent extinction, we also presented two original exemplars of both categories that were used during acquisition (4 trials in total); original exemplars from the CS+ category were reinforced with the pain-US. These 20 trials (8 GS+, 8 GS-, 2 CS+, and 2 CS-) were run in a random order. There were two important procedural differences with the acquisition phase: (a) on each trial, a startle probe was delivered during the GS (or CS) movement, but never during the ITI; and (b) on each trial, participants rated pain-US expectancy and how
afraid they were to perform the movement before actually carrying out the joystick movement.

**MAIN OUTCOME VARIABLES**

**Prospective Fear of Movement-Related Pain Ratings**
During the practice, acquisition, and generalization phases, participants rated how afraid they were to perform the “signaled” movements (CSs/GSs) before actually doing so. They were asked to rate “How fearful are you to perform this action?” on a 0 (Not fearful at all) to 10 (Very afraid) Likert scale.

**Prospective Pain-US Expectancy Ratings**
During the practice, acquisition, and generalization phases, participants rated the extent to which they expected the painful stimulus to occur when performing the “signaled” movements (CSs/GSs) on a 0 (Not at all) to 10 (Very much) Likert scale.

**Startle Eyeblink Modulation**
Orbicularis oculi electromyographic (EMG) activity was recorded with three Ag/AgCl Sensormedics electrodes (4 mm) filled with electrolyte gel. After cleaning the skin with exfoliating peeling cream to reduce interelectrode resistance, electrodes were placed on the left side of the face according to the site specifications proposed by Blumenthal et al. (2005). The raw signal was amplified by a Coulbourn isolated bioamplifier with bandpass filter (LabLinc v75–04). The recording bandwidth of the EMG signal was between 90 Hz and 1 kHz (±3 dB). The signal was rectified online and smoothed by a Coulbourn multifunction integrator (LabLinc v76–23 A) with a time constant of 20 ms. The EMG signal was digitized at 1,000 Hz from 200 ms before the onset of the auditory startle probe until 1,000 ms after probe onset.

**MANIPULATION CHECKS**

**Retrospective Pain-Related Fear**
As a manipulation check, after each conditioning block, we assessed how afraid participants were of the CS categories (opening vs. closing boxes). They answered the following question: “To what extent were you afraid to perform the action of opening/closing a box during the previous block?” on a Likert scale ranging from 0 (Not afraid at all) to 10 (Very afraid). To ensure that this assessment would not focus participants’ attention solely on the functional action categories that were of relevance to predict the pain, they also rated fear related to other perceptual and proprioceptive features that varied among the different exemplars of the functional categories but were not relevant to predict the pain: (a) movement direction, (b) size, and (c) color. Respectively, they answered the following questions on a Likert scale from 0 (Not afraid at all) to 10 (Very afraid): “To what extent were you afraid to perform the movement to the left/right during the previous block?”, “To what extent were you afraid of large/small boxes during the previous block?”, and “To what extent were you afraid of light/dark colored boxes during the previous block?”

**Pain-US Intensity and Unpleasantness**
After each conditioning block, participants answered the questions “How painful did you find the electrocutaneous stimulus in the previous block?” and “How unpleasant did you find the electrocutaneous stimulus in the previous block?” on 0 (Not at all) to 10 (Very much) Likert scales to monitor possible habituation or sensitization effects.

**EXPERIMENTAL SETTING**
Participants were seated in an armchair (0.6-meter screen distance) in a sound-attenuated and dimmed experimental room, adjacent to the experimenter’s room. Further verbal communication was possible through an intercom system; the experimenter observed the participants and their physiological responses online by means of a closed-circuit TV installation and computer monitors.

**RESPONSE DEFINITION AND DATA ANALYSIS OVERVIEW**

**Response Definition of the Startle Modulation**
Using PSPHA (Declercq, De Houwer, & Baeyens, 2008), a modular script-based program, we calculated the peak amplitudes defined as the maximum of the response curve within 21–175 ms after the startle probe onset. All startle waveforms were visually inspected off-line, and technical abnormalities and artifacts were eliminated using the PSPHA software. Every peak amplitude was scored by subtracting its baseline score (averaged EMG level between 1 and 20 ms after the probe onset). The raw scores were transformed to z scores to account for interindividual differences in physiological reactivity. In order to optimize the visualization of the startle data and avoid negative values on the y-axis, T scores—a linear transformation of the z scores—were used in the figures. Averages were calculated for responding during movements of CS/GS exemplars of both functional categories and ITI separately.

**Data Analysis Overview**
We carried out separate repeated measures (RM) ANOVAs on the respective dependent measures to examine the acquisition and generalization of pain-related fear based on conceptual knowledge. In particular, we hypothesized that (a) at the end of acquisition (ACQ4), fear and US-expectancy ratings...
as well as the eyeblink startle responses would be higher for the exemplars of the CS+ category than for the exemplars of the CS- category; and (b) during the test of generalization, fear and US-expectancy ratings as well as the eyeblink startle responses would be higher for novel exemplars of the CS+ category (GS+) than for novel exemplars of the CS- category (GS-). The $\alpha$ level was set at .05. In testing our a priori hypotheses, a Bonferroni correction was applied when using multiple planned comparisons. Greenhouse-Geisser corrections are reported when appropriate. Uncorrected degrees of freedom and corrected $p$ values are reported together with $\varepsilon$ and the effect size indication $\eta^2$ is reported for significant ANOVA effects and Cohen’s $d$ for planned comparisons. The interpretation thresholds based on Cohen (1988) and extended by Rosenthal (1996) can be summarized as follows: .20 = small, .50 = medium, .80 = large, and 1.30 = very large. Statistical analyses for all dependent measures were run with Statistica 12 software.

**Results**

**Prospective Fear of Movement-Related Pain Ratings**

**Practice**

We conducted a paired samples $t$ test on the mean pain-related fear ratings for the CS+ and CS- categories during the practice phase. As expected, before conditioning, fear ratings were low and did not differ between the CS+ and CS- categories, $t(49) = 1.59, p = .12$ (see Figure 2).

**Acquisition**

We carried out a 2 (Stimulus Category: CS+, CS-) $\times$ 4 (Block: ACQ1, ACQ2, ACQ3, ACQ4) RM ANOVA on the mean pain-related fear ratings for the CS categories during the four acquisition blocks (see Figure 2). There was a significant main effect of Stimulus Category, $F(1, 49) = 39.04, p < .0001, \eta^2 = .44$, and a significant main effect of Block, $F(3, 147) = 30.22, p < .0001, \varepsilon = .67, \eta^2 = .38$, both of which were qualified by a Stimulus Category $\times$ Block interaction, $F(3, 147) = 36.99, p < .0001, \varepsilon = .84, \eta^2 = .43$. Planned comparisons confirmed that at the end of the acquisition phase (ACQ4), the exemplars of the CS+ category elicited more pain-related fear than did those of the CS- category, $F(1, 49) = 36.45, p < .0001, d = 1.04$, although this was not the case in the beginning of the acquisition phase (ACQ1), $F < 1$.

**Test of Generalization**

We examined generalization of pain-related fear to the novel exemplars (GSs) of the learned CS categories, we ran an RM ANOVA (Stimulus Category: CS+, GS+, GS-, CS-; see Figure 2). There was a significant effect of Stimulus Category, $F(3, 147) = 38.83, \varepsilon = .41, p < .0001, \eta^2 = .44$. Planned comparisons confirmed that the original exemplars of the CS+ category still elicited more pain-related fear than did the original exemplars of the CS- category, $F(1, 49) = 36.45, p < .0001, d = .77$, suggesting that no extinction took place during the generalization test. More importantly and in line with our expectations,
participants reported more pain-related fear in response to the novel exemplars of the CS+ category (GS+) as compared with novel exemplars of the CS- category (GS-), \( F(1, 49) = 46.91, p < .0001, d = .86 \). No such differences occurred between the CS+ and the GS+ exemplars, \( F(1, 49) = 2.60, p = .11, d = .07 \), or the CS- and the GS- exemplars, \( F(1, 49) = 1.36, p = .25, d = .05 \), suggesting that there was a perfect transfer of the learned contingencies based on a certain set of exemplars to novel exemplars of the same functional category.

PROSPECTIVE PAIN-US EXPECTANCY RATINGs

**Practice**
We conducted a paired samples \( t \) test on the mean pain-US expectancy ratings for the CS+ and CS- categories during the practice phase. As expected, before conditioning, pain-US expectancy ratings were low and did not differ between the CS+ and CS- categories, \( t(49) = 0.41, p = .68 \) (see Figure 3).

**Acquisition**
We carried out a 2 (Stimulus Category: CS+, CS-) \( \times \) 4 (Block: ACQ1, ACQ2, ACQ3, ACQ4) RM ANOVA on the mean pain-US expectancy ratings for the CS categories during the four acquisition blocks (see Figure 3). There was a significant main effect of Stimulus Category, \( F(1, 49) = 62.02, p < .0001, \eta^2_p = .56 \), but not of Block \( (F < 1) \). This was qualified by a significant Stimulus Category \( \times \) Block interaction, \( F(3, 147) = 45.80, p < .0001, \epsilon = .84, \eta^2_p = .48 \). Planned comparisons confirmed that at the end of the acquisition phase (ACQ4), the CS+ category induced higher pain-US expectancies than did the CS- category, \( F(1, 49) = 78.91, p < .0001, d = 1.59 \), although this was not the case in the beginning of the acquisition phase (ACQ1), \( F < 1 \).

**Test of Generalization**
To examine generalization of pain-US expectancy to the novel exemplars (GSs) of the learned CS categories, we ran a RM ANOVA (Stimulus Category: CS+, GS+, GS-, CS-) (see Figure 3). There was a significant effect of Stimulus Category, \( F(3, 147) = 42.47, \epsilon = .47, p < .0001, \eta^2_p = .46 \). Planned comparisons confirmed that participants still expected the pain-US to occur more after the original exemplars of the CS+ category than the original exemplars of the CS- category, \( F(1, 49) = 32.35, p < .0001, d = .88 \). More importantly and in line with the pain-related fear ratings, participants showed higher pain-US expectancies in response to the novel exemplars of the CS+ category (GS+) as compared with novel exemplars of the CS- category (GS-), \( F(1, 49) = 66.74, p < .0001, d = 1.25 \). No such differences occurred between the CS+ and the GS+ exemplars, \( F(1, 49) = 2.91, p = .09, d = .13 \), or the CS- and the GS- exemplars, \( F < 1 \).

STARTLE EYEBLINK MODULATION

**Acquisition**
We carried out a 3 (Stimulus Type: CS+, CS-, ITI) \( \times \) 4 (Block: ACQ1, ACQ2, ACQ3, ACQ4) RM ANOVA on the mean startle amplitudes of the CS+ and CS- categories as well as the ITI during the acquisition phase (see Figure 4). This analysis yielded a significant
main effect of Block, $F(3, 147) = 19.74$, $\epsilon = .58$, $p < .0001$, $\eta^2_p = .29$, and a significant main effect of Stimulus Type, $F(2, 98) = 76.93$, $\epsilon = .81$, $p < .0001$, $\eta^2_p = .61$, both of which were qualified by a Stimulus Type × Block interaction, $F(6, 294) = 3.32$, $\epsilon = .77$, $p < .01$, $\eta^2_p = .06$. Although startle amplitudes during the exemplars of the CS+ category tended to be elevated compared with those of the CS- category, planned comparisons did not support statistical significance, $F(1, 49) = 3.25$, $p = .08$, $d = .25$. Interestingly, startle amplitudes during both CS categories were significantly higher than during the ITI, $F(1, 49) = 105.66$, $p < .0001$, $d = 1.11$, suggesting that both CS categories showed elevated psychophysiological responding as compared with the safe context.

**Test of Generalization**

Because there were no reliable acquisition effects in the startle eyeblink measures, generalization effects were not further reported.

**Manipulation Checks**

**Retrospective Pain-Related Fear**

To ensure that the other irrelevant features of the exemplars of the CS+ category elicited less fear than did the relevant feature (i.e., the function of opening/closing a box), we conducted a 5 (Block: ACQ1, ACQ2, ACQ3, ACQ4, Test) × 8 (Stimulus Category: CS+, CS-, left, right, dark, light, big, small) RM ANOVA on retrospective pain-related fear ratings (see Figure 5). This analysis showed a significant main effect of Block, $F(4, 196) = 9.58$, $\epsilon = .80$, $p < .0001$, $\eta^2_p = .16$, suggesting that over time, ratings of both unpleasantness and intensity changed. There was also a significant main effect of Rating, $F(1, 49) = 39.11$, $p < .0001$, $\eta^2_p = .44$, indicating that unpleasantness ratings were higher than pain-US intensity ratings. The Rating × Block interaction was not significant, $F(4, 196) = 1.66$, $\epsilon = .80$, $p = .17$, $\eta^2_p = .03$, suggesting that the pain-US intensity and unpleasantness ratings demonstrated a similar pattern of change. In particular, there was a slight increase in pain-US intensity (linear trend;
Previous research has shown that perceptual/proprioceptive similarity with a movement that featured in a painful episode facilitates the generalization of conditioned pain-related fear to novel movements (Meulders et al., 2013; Meulders & Vlaeyen, 2013). This type of fear generalization has been referred to as perceptual pain-related fear generalization. A hallmark of human cognition is the ability to additionally extract conceptual knowledge from a learning episode; however, research on this conceptual pathway to fear generalization is scant. The aim of the current study was to investigate the acquisition and generalization of pain-related fear based on conceptual knowledge relating to category membership.

First, we successfully established the acquisition of fear of movement-related pain based on a superordinate “action” category membership. In particular, participants learned that unique exemplars of one action category (CS+; e.g., opening boxes) predicted a painful electrocutaneous stimulus (pain-US), whereas unique exemplars of another action category (CS-; e.g., closing boxes) did not. This pattern was observed in self-reported fear of movement-related pain, as well as in the pain-US expectancy ratings. In contrast, eyeblink startle responses were not elevated in response to the exemplars of the CS+ category as compared with those of the CS- category, but exemplars of both CS+ and CS- action categories elicited higher startle responses than did the context alone (i.e., ITI startle responses).

Second, we demonstrated the spreading of conditioned responding to novel exemplars of the learned CS+ category, but not to the CS- category.
Participants not only reported more pain-related fear in response to the novel exemplars of the CS+ category (GS+) as compared with novel exemplars of the CS- category (GS-), but also showed higher pain-US expectancies to the GS+ than to the GS-. In other words, participants reported being more afraid and expected the pain to occur more when they had to open novel boxes (with different combinations of size, color, and shape than the original boxes that were used during acquisition) as compared with when they had to close novel boxes, or vice versa depending on which action (opening/closing boxes) was paired with pain during acquisition. Because there was no reliable acquisition effect in the startle eyeblink measures, no proper test of generalization could be performed. This is in contrast with an earlier study on perceptual pain-related fear generalization by Meulders et al. (2013), who showed a perceptual generalization gradient in the startle eyeblink measures, but failed to observe a similar gradient in the self-reported fear measures. This discrepancy might be explained by several methodological differences between the current study and that of Meulders et al. (2013): (a) Meulders et al. (2013) tested generalization under extinction (i.e., original CSs were not reinforced), which typically leads to quick decline of fear responding; (b) because self-reported fear was not assessed prospectively but in retrospect, fear of movement-related pain might have already extinguished; and (c) because startle eyeblink measures were collected during the generalization trials, they were less affected by extinction (especially in the first trials). Closely related to the current findings, previous research by Bennett, Meulders, Baeyens, and Vlaeyen (2015) has shown that neutral movements’ conceptual relationship with pain-associated stimuli can precipitate fear of movement-related pain. In their experiment, two artificial categories were created so that neutral joystick arm movements and nonsense words were equivalent. During the subsequent pain-related fear conditioning procedure, nonsense words of one category were paired with a pain-US (B+), whereas nonsense words from another category were followed by no pain-US (B-). During the symbolic generalization test, participants were asked to perform specific joystick movements that were trained to be either conceptually similar (C+) or dissimilar (C-) to the pain-associated nonsense words. Results showed that conceptual equivalence between neutral movements and pain-associated nonsense words (C+) evoked elevated pain-related fear, pain-US expectancy, and unpleasantness compared with movements conceptually related to nonsense words that were not associated with pain (C-). Like the novel exemplars of the CS+ category in the current study, the movements themselves were never paired with a pain-US, nor were the movements in any sense perceptually similar to the pain-associated nonsense words. The present findings further corroborate the idea that category membership based on derived equivalence can facilitate the spreading of pain-related fear.

Some remarkable findings deserve further attention. First, we observed a discrepancy between the self-reported and psychophysiological fear measures. Dissociations between response systems have been reported before (Soeter & Kindt, 2010) and do not necessarily reflect a failure to replicate. It has been argued that dissociations between declarative knowledge (CS-US contingency awareness) and the psychophysiological fear response might also pertain to relevant information, as both measures might tap into different aspects of fear (fear is associated with both expectancy of harm and elevated psychophysiological arousal; Lang, Bradley, & Cuthbert, 1998). For example, due to its salience, visual information is often prioritized and captures attention more quickly than other sensory information (tactile, auditory, or proprioceptive information). Because other “irrelevant,” mostly visual features such as the size, color, and shape of the boxes, as well as the direction in which participants had to move the joystick, overlapped between CS+ and CS- categories, a possible explanation may be that these features are processed earlier than the relevant “superordinate” action category information. Consequently, these stimulus features might elicit a certain level of fearful arousal leading to nondifferential facilitation of startle responses during exemplars of both CS+ and CS- categories compared with ITI startle responses (i.e., contextual fear). Partial evidence supporting this post hoc explanation is found in the retrospective pain-related fear ratings showing that next to the expected action category-based differential fear learning (e.g., opening vs. closing boxes), the irrelevant categories such as movement direction, size, and color brightness of the boxes generated fear reports situated between the CS+ and CS- categories responses.

Another explanation might be that extracting conceptual information (i.e., categorizing novel exemplars) is a cognitively demanding task. Previous studies have reported that cognitive load can inhibit the startle response (Acocella & Blumenthal, 1990; Filion, Dawson, & Schell, 1993; Schell, Wynn, Dawson, Sinai, & Niebala, 2000), yet the observed data pattern does not seem to support this explanation because startle responses to both CS+ and CS- categories were enhanced compared with ITI startle responses. Alternatively, it seems feasible that information mediated by higher-order cognition such as category information also requires more
time and effort to process, which in turn raises the possibility that short-latency reflexes such as the eyeblink startle response may not be inhibited by safety cues or facilitated by danger signals based on effortful categorization. On a related note, it is also possible that the startle probes were presented too early after action onset for participants to classify the action as belonging to the CS+ or CS- categories and to produce startle modulation effects. To our knowledge, this is the first study employing startle eyeblink measures in the context of category-based fear conditioning. In fact, at least three important features differ between our study and that of Dunsmoor et al. (2012): CS modality, CS duration, and the type of psychophysiological measure used. In Dunsmoor et al. (2012), visual stimuli presented for 6 seconds served as CSs, and skin conductance responding (SCR) was used as a psychophysiological index of category-based fear learning. The SCR is a longer-latency autonomic response typically emerging around 1 second after stimulus onset and peaking a maximum of 5 seconds later (for visual stimuli), whereas the startle response typically starts 20–50 ms after the startle-eliciting stimulus. This means that the longer stimulus duration (which was possible due to the different modality—i.e., visual stimuli) might have allowed participants to categorize the stimuli on time to generate category-specific anticipatory SCRs. To settle these unresolved issues regarding the observed discrepancy between self-reports and psychophysiological measures, future research may include SCR measures, postpone the presentation of the startle probe after action onset, and preclude the overlap of irrelevant features such as size, shape, and color of the boxes in the CS+ and CS- categories in order to create mutual exclusivity.

Second, compared with previous studies investigating fear of movement-related pain using the VJM paradigm (Meulders et al., 2011; Meulders & Vlaeyen, 2013), differential category-based fear of movement-related pain took longer to develop. Acquisition may have been slowed down because participants had to extract the relevant information relating to the occurrence of the pain-US for each single exemplar and then had to create categories based on this abstract knowledge; in that sense, acquisition already involved a kind of generalization from each single exemplar to the category. We also found that, once acquired, category-based fear of movement-related pain transferred rigorously to novel exemplars of the CS+ category, although these generalization stimuli were tested under extinction (i.e., GSs were never reinforced, two out of four of the original CS+ exemplars were reinforced). Compared with previous studies investigating the extinction of fear of movement-related pain using the VJM paradigm (Meulders & Vlaeyen, 2012), extinction learning appears to be slowed down as well. Further research should include more extinction trials to define the boundaries for category-based pain-related fear extinction.

Some limitations of our study should be addressed. First, for practical reasons, we chose to use overlapping exemplar features (e.g., size, color, shape, movement direction) in both of the CS categories. This might have caused the nondifferential elevation of startle responses. Future research should attempt to disentangle whether there is a genuine discrepancy between self-reports and startle responses or whether this is an artifact of the present procedure. Second, in contrast with Dunsmoor et al. (2012), and also for practical reasons, we presented the same exemplars in each conditioning block during acquisition. Therefore, it is possible that acquisition would be even more slowed down when unique exemplars were presented during each acquisition block. Third, there might be a minimal perceptual similarity between the different exemplars of “opening boxes” and “closing boxes” because for each exemplar in the closing boxes category, there was a lid that was sticking out (although the position, shape, size, and angle differed), whereas for the opening boxes category the lid was not visible (i.e., closed box; see Appendix 1). Future research may employ more distinct exemplars of the action categories in order to minimize this low-level perceptual overlap. For example, GSs such as closing/opening doors or jars may be a good method to take the test of category-based generalization a step further. Fourth, to study the basic learning mechanisms involved in category-based pain-related fear generalization in the lab, we used a sample of relatively young healthy subjects without any psychiatric disorders or medical illnesses. Given that comorbid anxiety, depression, and other medical illnesses are common in people with chronic pain, these learning processes might play out differently in clinical samples. Therefore, future research is needed to validate these findings in chronic pain populations. Finally, our main findings in the self-reported fear and US-expectancy were not replicated in the eyeblink startle measures. Cognitive models of anxiety propose that (conditioned) fear is closely

1 On a related note, in the study of Dunsmoor et al. (2012), the influence of perceptual features also could not be fully ruled out. In their study, unique exemplars of the superordinate category “animals” served as the CS+ and exemplars of the superordinate category “tools” served as the CS-, or vice versa. It can be argued that pictures of tools typically comprise more straight lines than pictures of animals, which might render the exemplars within one category also perceptually more similar.
related to the formation of threat beliefs and harm expectation (Chan & Lovibond, 1996). Yet, some researchers have argued that verbal measures such as US-expectancy lack objectivity, as they may be susceptible to effects of social desirability and experimental demand (Craske, Hermans, & Vansteenwegen, 2006). However, it has been argued that (a) US-expectancy measures have face validity, construct validity, predictive validity, and diagnostic validity with respect to anxiety-related disorders (Boddez, Baeyens, Luyten, et al., 2013a); (b) in ambiguous and complex fear-conditioning procedures such as this one, demand effects are not a large concern because the desired responses and study hypotheses are not obvious (Boddez, Baeyens, Hermans, & Beckers, 2013b); (c) such complex designs may not yield differential effects on psychophysiological indices because these measures are not always sufficiently sensitive to detect subtle differences in modulation between multiple stimuli (Ahmed & Lovibond, 2015); and (d) modulation of the startle eyeblink is not a unique measure for fear, as sometimes claimed. For example, it is also modulated by attention (Lipp, Cox, & Siddle, 2001), which might have been an issue in the current study because unique exemplars of the CS categories were used within each block. As a consequence, the orienting response toward all relatively new stimuli might explain the nondifferential potentiation of the startle response.

We have argued that fear learning and generalization is adaptive to a certain degree. The crucial difference between adaptive fear and pathological fear is often conceptualized in terms of fear intensity and avoidance behavior (Dymond et al., 2015). In the case of chronic musculoskeletal pain, fear of movement-related pain becomes pathological to the extent that it impairs normal functioning and prompts avoidance of such movements. Although intensity and avoidance are important in the transition from adaptive to maladaptive fear, we believe that fear overgeneralization, or the spreading of fear to technically safe movements, may be of particular relevance. Although surprisingly few studies addressed fear generalization in the field of pain-related fear, some findings may be of clinical importance.

First and most importantly, treatments that target pain-related fear in chronic pain (e.g., exposure-based therapy) require an in-depth appraisal of the crucial stimuli and their learning history (den Hollander et al., 2016; Vlaeyen, de Jong, Geilen, Heuts, & van Breukelen, 2001). An important conclusion from generalization research is that fear can be triggered by stimuli that never featured in a pain episode. The present study indeed shows that fear can be evoked by stimuli that are perceptually not related to the original CSs and that were never associated with pain themselves. Therefore, the notion of conceptual fear generalization might shed light on the fact that conditioning experiences are often difficult to reconstruct in a clinical context (Poulton & Menzies, 2002). However, a more comprehensive understanding of nonperceptual fear generalization is needed before clear recommendations can be given about how these learning processes may impact the conventional treatment approach.

Second, given that generalization may be crucial in the transition from normal to pathological fear, it might be targeted in the prevention of chronic pain disability. For example, discrimination training before a painful medical procedure may be a promising pathway to prevent the spreading of fear from movements and situations that should be avoided to other perceptually or conceptually related events that do not need to be feared or avoided (see Vervliet, Kindt, Vansteenwegen, & Hermans, 2010, for a similar argument).

To conclude, we demonstrated that pain-related fear can be acquired based on conceptual knowledge, and that this concept-based fear of movement-related pain generalizes to novel exemplars of the CS+ category but not to those of the CS- category. We believe that these novel developments in fear generalization research focusing on the integration with higher-order reasoning and conceptual knowledge are crucial to capture the interplay of pathways involved in the spreading of fear and avoidance behavior in chronic pain conditions.

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Conflict of Interest Statement
The authors declare that there are no conflicts of interest.
References


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