Nonconscious activation of placebo and nocebo pain responses

Karin B. Jensen\textsuperscript{a,b,c,1}, Ted J. Kaptchuk\textsuperscript{b}, Irving Kirsch\textsuperscript{b,d}, Jacqueline Raicek\textsuperscript{b}, Kara M. Lindstrom\textsuperscript{b}, Chantal Berna\textsuperscript{b,f}, Randy L. Gollub\textsuperscript{b,c,1}, Martin Ingvar\textsuperscript{a}, and Jian Kong\textsuperscript{a,b,c}

\textsuperscript{a}Department of Psychiatry, Massachusetts General Hospital/Harvard Medical School, Charlestown, MA 02129; \textsuperscript{b}Program in Placebo Studies, Beth Israel Deaconess Medical Center/Harvard Medical School, Boston, MA 02215; \textsuperscript{c}MGH/MIT/HMS Athinoula A. Martinos Center for Biomedical Imaging, Charlestown, MA; \textsuperscript{d}School of Psychology, Plymouth University, Plymouth PL4 8AA, United Kingdom; \textsuperscript{e}Department of Clinical Neuroscience, Osher Center for Integrative Medicine, Karolinska Institutet, 17176 Stockholm, Sweden; and \textsuperscript{f}Division of Pain Medicine, Department of Anesthesia, Critical Care and Pain Medicine, Massachusetts General Hospital, Boston, MA 02108

Edited by Tomas G. M. Hökfelt, Karolinska Institutet, Stockholm, Sweden, and approved August 6, 2012 (received for review February 3, 2012)

The dominant theories of human placebo effects rely on a notion that consciously perceptible cues, such as verbal information or distinct stimuli in classical conditioning, provide signals that activate placebo effects. However, growing evidence suggest that behavior can be triggered by stimuli presented outside of conscious awareness. Here, we performed two experiments in which the responses to thermal pain stimuli were assessed. The first experiment assessed whether a conditioning paradigm, using clearly visible cues for high and low pain, could induce placebo and nocebo responses. The second experiment, in a separate group of subjects, assessed whether conditioned placebo and nocebo responses could be triggered in response to nonconscious (masked) exposures to the same cues. A total of 40 healthy volunteers (24 female, mean age 23 y) were investigated in a laboratory setting. Participants rated each pain stimulus on a numeric response scale, ranging from 0 = no pain to 100 = worst imaginable pain. Significant placebo and nocebo effects were found in both experiment 1 (using clearly visible stimuli) and experiment 2 (using nonconscious stimuli), indicating that the mechanisms responsible for placebo and nocebo effects can operate without conscious awareness of the triggering cues. This is a unique experimental verification of the influence of nonconscious conditioned stimuli on placebo/nocebo effects and the results challenge the exclusive role of awareness and conscious cognitions in placebo responses.

analgesia | hyperalgesia | consciousness

Placebo and nocebo effects are critical components of medical practice and clinical research. Placebo analgesia and nocebo hyperalgesia are the most robust and well studied of these effects. Learning is known to play an important role in placebo and nocebo effects and the dominant theories invoke classical conditioning and expectancies as explanatory tools (1). Both rely on a notion that the conscious perception of sensory or social stimuli, such as the cue that triggers expectation or the conditioned stimulus in classical conditioning, are needed to obtain placebo responses. In some circumstances, conditioning may be an automatic nonconscious process, but in most cases, it seems to involve the formation of expectations (2–4). However, it is not known whether conscious perception of a conditioned stimulus is needed to elicit a conditioned response.

There is a large literature suggesting that behavior can be motivated by stimuli that are not consciously perceived, because they are presented at low intensities or masked from conscious awareness (5, 6), sometimes referred to as subliminal stimuli. Nonconscious operations are considered a fundamental feature of human cognition, for example in reward processing (7, 8), fear learning (9, 10), and social behavior (11, 12). Furthermore, evidence suggests that conditioned responses may be acquired outside of conscious awareness (13–15). Neuroimaging studies of the human brain suggest that certain structures, such as the striatum and the amygdala, can process incoming stimuli before they reach conscious awareness, and thus they may mediate nonconscious effects on human cognition and behavior (16, 17). It has never been investigated, however, whether learned placebo and nocebo responses can be triggered through this cerebral circuit that bypasses conscious awareness.

Placebo and nocebo may be seen as the behavioral response to signals of reward and threat, respectively. Considering the neurobiological evidence for nonconscious processing of reward and threat signals, placebo and nocebo responses would have the potential to be activated by masked stimuli. Here, we experimentally test this hypothesis and explore the role of nonconscious mental processes in placebo analgesia by investigating whether conditioned placebo and nocebo responses can be activated by masked stimuli ($n = 40$). Conditioning was performed with clearly visible cues and a subsequent test-sequence–measured placebo and nocebo responses to either visible (unmasked) or nonconscious (masked) cues (see Fig. 1). Our goal was to test whether conditioned placebo and nocebo responses could be activated by both consciously and nonconsciously perceived cues.

Results

Experiment 1. Experiment 1 was designed to ascertain that conditioned placebo and nocebo pain responses could beelicited by consciously perceived stimuli. An initial conditioning sequence, in which high and low thermal pain stimuli were paired with clearly visible exposures of two male faces on a computer screen, produced a mean rating of “high pain” at 63 ($\pm 22.1$) on a numeric response scale (NRS), ranging from 0 (no pain) to 100 (worst imaginable pain), and “low pain” at 24 ($\pm 14.7$) on the NRS. The test sequence, also using clearly visible stimuli, revealed significantly higher pain ratings in response to the high face and lower ratings for the low face, despite identical moderate temperatures. The control condition, paired with the same moderate temperature, resulted in pain ratings in between the high and low conditions; ANOVA main effect for face type (high/low/control) $F(2, 36) = 24, P < 0.001$. All pairwise comparisons between the high/low ($P < 0.001$), high/control ($P < 0.001$) and low/control ($P = 0.003$) conditions were significant (Fig. 2) and all subjects reported that they could clearly discriminate between the different faces during the test sequence. Correlations (Pearson’s $r$) revealed that there was a positive correlation between the difference in high minus low pain ratings during conditioning and the high minus low pain rating during the test sequence of experiment 1; $r = 0.608, P = 0.006$ (Fig. 3). Analyses of alpha reliability (Chronbach’s alpha) showed that the three conditions of


The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

\textsuperscript{1}To whom correspondence should be addressed. E-mail: karinj@nmr.mgh.harvard.edu.

www.pnas.org/cgi/doi/10.1073/pnas.1202056109

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the test sequence in experiment 1 (high face/low face/control face) displayed a reliability of 0.98. The pain ratings during the test sequence were not affected by the factor “time” \(F(2, 32) = 2.8, P = 0.073\) and there was no interaction between pain ratings × time \(F(2, 32) = 2.2, P = 0.134\), validating that pain habituation or sensitization did not confound the placebo/nocebo responses.

In conclusion, the results from experiment 1 showed significant placebo and nocebo responses and also indicated that their magnitude was predicted by the difference between high and low pain ratings during the learning phase.

**Experiment 2.** Experiment 2 was designed to test the unique hypothesis that conditioned placebo and nocebo responses could be triggered by masked, nonconscious stimuli. In most respects,
the methods in experiment 2 were identical to those used in experiment 1. The only difference was that all faces during the test sequence were presented for 12 ms followed by a visual mask for 84 ms, resulting in comparable visual exposures for both experiments (faces shown for 100 ms in experiment 1). The conditioning sequence in experiment 2 was performed with clearly recognizable images and produced a mean rating of “high pain” at 52 (± 19) and “low pain” at 21 (± 11.8) NRS. The test sequence, however, was performed using masked images, preventing subjects from consciously recognizing the stimuli. As in experiment 1, the results from the test sequence revealed significantly higher pain ratings in response to the high face, lower ratings for the low face, and intermediate ratings for the control face, despite identical moderate temperatures, ANOVA main effect for face type: F(1, 18) = 11, P < 0.001. All pairwise comparisons in experiment 2 were also significant despite subjects’ inability to consciously distinguish the faces from each other; high/low (P = 0.002), high/control (P = 0.011), and low/control (P = 0.008) (Fig. 2). A correlation analysis (Pearson’s) revealed a positive correlation between the difference in high minus low pain ratings during conditioning and the high minus low pain rating during the test sequence (r = 0.822, P < 0.001), indicating that the magnitude of masked placebo/nocebo responses could also be predicted by pain ratings during conditioning. Analyses of alpha reliability (Chronbach’s alpha) showed that the three conditions of the test sequence in experiment 2 (masked high face/low face/control face) displayed a reliability of 0.97. The pain ratings during the test sequence of experiment 2 were not affected by the factor time F(2, 38) = 0.8, P = 0.449 and there was no interaction between pain ratings x time F(2, 32) = 2.2, P = 0.134, validating that pain habituation or sensitization did not confound the placebo/nocebo responses.

All subjects in experiment 2 reported being unable to consciously discriminate between the different faces during the test sequence. To control for any potential variance due to differences in recognition of the faces in experiment 2, the outcome from a subsequent masked face recognition test was used as a covariate in the overall ANOVA, which demonstrated nonsignificance, F(1, 18) = 0.69, P = 0.414. Further validation that the images were indeed nonconscious was demonstrated by an analysis revealing that subjects’ ability to recognize the masked images, represented by the individual recognition rates, was uncorrelated to the effect of high minus low pain during the test sequence (r = −0.253, P = 0.296).

**Discussion**

Results from the present study demonstrate that placebo and nocebo mechanisms can be triggered by nonconscious cues, operating outside of conscious awareness. As far back as 1885, Peirce and Jastrow suggested that nonconscious cues could influence somatosensory perception (18). Since then, a large literature has suggested that nonconscious signals of threat and relief can be processed by subcortical (but also cortical) structures in the brain and influence behavioral outcomes (8, 19). Recent theoretical analyses have also suggested the possibility that placebo and nocebo responses may be mediated by nonconscious operations (20, 21), what Kihlstrom called the “cognitive unconscious” (22). Previous studies (13–15) have demonstrated that associative learning can be obtained by the use of nonconscious stimuli during the acquisition of conditioned responses. In the present study, we extended the understanding of nonconscious cognitions by showing that explicitly conditioned placebo analgesia and nocebo hyperalgesic responses can be activated by nonconscious cues. Our results thereby translate the investigation of nonconscious effects to the clinical realm, by suggesting that health-related responses can be triggered by cues that are not consciously perceived, not only for pain, which is one of the most common reasons for seeking healthcare (23), but also for other medical problems with demonstrated placebo effects, e.g., asthma (24), depression (25), and irritable bowel syndrome (26). Understanding the role of nonconscious processes in placebo/nocebo opens unique possibilities of enhancing clinical care by attending
to the impact of nonconscious cues conveyed during the therapeu
tic encounter and improving therapeutic decisions.

Common theories of placebos involve expectancy and classical conditioning, and both mechanisms, although they can be hard to separate, involve conscious perception of the stimulus that elicits the placebo or nocebo response (3, 4). Our study is clearly dis
tinguished from previous studies because it focuses on the nonconscious activation of placebo/nocebo responses and demon
strates that placebo/nocebo can be activated even if the con
tioned stimulus is not consciously perceived. In traditional placebo studies, conditioning is often used by pairing the ad
ministration of an unconditioned stimulus (e.g., effective anal
gesic pill, cream, or injection) with a conditioned stimulus (e.g.,
inert placebo pill, cream, or injection), thus producing placebo responses through “associative learning” (27–30). Even if asso
ciative learning in such studies has been described in terms of a “nonconscious” mechanism (in which cognition may be an
epiphemomenon, rather than part of a causal chain), the activa
tion of the conditioned response has always been obtained by
a perceptible conditioned stimulus. Furthermore, much of the
evidence for conditioning effects in human placebo experiments
demonstrates that a conscious cognitive component plays a sig
ificant role in the placebo conditioning. For example, Mont
gomery and Kirsch (31) replicated Voudouris et al.’s (32, 33)
early conditioning placebo experiments and found that conscious
awareness of the conditioning process eradicated placebo con
ditioning. Recent work by Watson et al. (34, 35) also supports
the notion that conscious expectation plays a dominant role in
conditioned placebo responses. On the other hand, in one of the
most compelling experiments of placebo effects and conditioning,
Benedetti and colleagues (36) showed that conditioned im
mune and endocrine placebo responses could still be elicited by
saline placebo even if the subjects were given clear verbal
directions not to expect any positive change. Given that subjects
were aware of the injections of saline (the conditioned stimu
i) in this experiment, Benedetti later noted that “a [remaining] key
question is: does unconscious conditioning exist in humans?”
(ref. 37, p. 53). To the best of our knowledge, our study presents
unique evidence that conditioned placebo responses can be ac
tivated by cues outside of conscious awareness.

Our results and proposed model, shed light on findings from
two previous placebo/nocebo studies that investigated how pain
ratings can be affected by the interaction between patients’ and
clinicians’ expectancies of pain relief. In one study, Gracely et al.
(38) found, in a double-blind experiment of pain relief, that the
clinician’s a priori knowledge of the likelihood of administering
active analgesic treatment versus placebo was transmitted to the
patient and influenced the placebo response. In another study,
Levine and Gordon (39) compared the double-blind administra
tion of morphine/placebo by either a hidden person or a hid
den machine and found that the placebo response was sig
ificantly lower in response to the machine. We speculate that
in both studies, subtle cues that the clinician conveyed to the
patient may have been perceived without conscious awareness.

The outcomes from these experiments suggest that placebo and
nocebo effects in the clinical setting might not only be induced
through explicit instructions and explanations, but also through
nonconscious cues embedded in the patient–clinician in
teraction. Nevertheless, neither of these experiments explicitly
tested the hypothesis that the findings were due to nonconscious
cue mechanisms.

The present study demonstrated successful activation of pla
cebo and nocebo effects in responses to both explicit (experiment 1) and nonconscious cues (experiment 2), suggesting that
different levels of brain processing may be involved. Previous
placebo and nocebo neuroimaging studies (40–44), using explicit
cues to evoke placebo responses, conclude that conscious pla
cebo effects recruit a combination of cortical and subcortical
brain regions to modulate pain. Brain imaging studies also sug
ggest that the brain can process environmental cues even if they
are not reaching conscious awareness, largely through sub
cortical regions of the brain such as the amygdala and ventral striatum (8, 10). Thus, we speculate that nonconscious cues may
work through the subcortical regions of the brain to produce
placebo and nocebo effects. Furthermore, we speculate that
nonconscious placebo and nocebo effects may not use the
commonly reported cortical regions of the brain, such as the
rostral anterior cingulate cortex (40, 43) and the prefrontal
cortex (41, 42). Conversely, they are likely to be processed in
subcortical parts of the brain that rely on a minimal account of
awareness, such as the basal ganglia.

In summary, the present study provides an experimental
demonstration of the influence of consciously nonrecognized
stimuli on conditioned placebo and nocebo responses. It suggests
that cognitive modulation of pain can be exerted without con
scious awareness of the triggering cues. Our results point to the
importance of a care process where the trajectory toward health
is seen as a learning experience that is highly influenced by
the activation of nonconscious environmental cues. Future studies
will show whether the present findings can be translated into
a clinical setting where nonconscious effects on health-related
behavior and treatment outcomes can be further validated. In
addition to pain responses, conditioned placebo effects are
known to affect a variety of clinical symptoms (24, 45–47), sug
ggesting that the present findings could be translated to other
disciplines than pain. In addition, future studies should establish
whether conditioned placebo and nocebo responses could be
obtained with the use of nonconscious stimuli also during the
acquisition phase of the conditioning procedure.

Materials and Methods

In total, 40 healthy subjects were included in this study (experiment 1: n = 20
subjects, 13 women and 7 men, mean age 22 ± 4 and experiment 2: n = 20;
11 women and 9 men, mean age 24 ± 4). All subjects were right handed,
with no history of medical or psychiatric illness and no previous experience
with fast image exposures or backward-masking experiments. Subjects were
recruited through posted flyers at several different universities and at free
expression boards in residential buildings.

Thermal pain stimuli were delivered using the Pathway CHEPS system from
Medoc, with a 27-mm diameter CHEPS thermode. The calibrated goal tem
peratures were reached with a ramp up time of 300 ms and the duration of
each pain stimulus was 3 s. An 85-Hz, 17-inch cathode ray tube monitor (NEC
AccuSync) was used for visual presentations and the masked stimulus pre
sentations were synchronized with the pain onset (1 ms). Presentation resolu
tion was 1,024 × 768 pixels and the experiment was programmed in
Presentation 13.0 (Neurobehavioral Systems). The images used in the current
experiment were taken from The Karolinska Directed Emotional Faces set
(KDEF) (48), a set of images specifically developed for use in perception,
attention, emotion, memory, and backward masking experiments. The whole
set consists of 70 individuals (35 male, 35 female), mean age 25 y (range
20–30) with seven different facial expressions per individual. The images
used in the present experiment only represented men in control
expressions, i.e., no emotional valence. In total, 24 male faces were used for
the purpose of this study.

Subjects were screened for inclusion and exclusion criteria over the tele
phone and then scheduled for an experiment. Subjects were informed that
the study investigated “the influence of implicit and explicit learning on pain
perception” but the full purpose of the study was not revealed until the
experiment was over. All subjects gave written informed consent and the
study was approved by The Institutional Review Board at Massachusetts
General Hospital.

The experiment was carried out in a quiet room at constant temperature
(23 °C). Subjects were seated in front of a desk with the monitor placed
straight in front of them: −70 cm from the subject’s face. The desk faced
a wall, preventing subjects from visual distractions. The thermode stimulator
was placed on the subject’s left volar forearm. Calibration of high and low
pain was performed by means of ascending temperatures, starting from
40 °C, followed by a randomized series of mixed high and low temperatures.
The goal was to find temperatures that would elicit high pain at −60 of 100
on a 0–100 NRS and low pain at ~20 NRS. The difference between the chosen high and low pain temperature was fixed to 3 °C for all subjects, e.g., high pain–low pain would be represented by 49°/46 °C in one individual and 47°/44 °C in another.

After subjective calibration of heat pain, subjects were given the following instruction for the conditioning sequence: “You are about to see some pictures on the screen. Each picture is paired with a pain stimulus on your arm. Your task is to focus on the screen at all times and after each picture you would like to rate how much pain you felt on your arm, using the same 0–100 verbal scale that you used during the calibration.” To ensure that subjects maintained high attention, the conditioning sequence was divided in two blocks of ~7 min each. In between the two blocks, subjects had the chance to stretch their legs and look away from the monitor for about 1.5 min or as long as they needed. In total, 50 stimuli were presented during the conditioning sequence: 25 for the high-pain face and 25 for the low-pain face. The exposure time of each image was 100 ms and the mean stimulus onset asynchrony (SOA) was 15 s (range 13–17 s). In both experiment 1 and experiment 2, the two male faces associated with high or low pain were counterbalanced to reduce the possibility that a certain face would contribute to high or low pain ratings. Immediately after the conditioning sequence, subjects were given the following instruction: “You are about to see the same pictures on the screen again and each picture will be paired with a pain stimulus on your arm, just like before. The only difference is that this time there will also be pictures of new guys, that you haven’t been exposed to before. Your task is to focus on the screen at all times and after each picture I would like you to rate how much pain you felt on your arm using the 0–100 verbal scale.” In experiment 2, the following sentence was also added: “During this sequence, the pictures will be shown to you much faster than before, and you might not be able to recognize them. This is normal and something that we programmed on purpose. Your only task is to focus on the screen at all times and rate the pain on your arm, even if you can’t see the pictures.” The test sequence consisted of 60 stimuli: 20 for the high-pain condition, 20 for the low-pain condition, and 20 for the control condition. The test sequence was divided in three 6-min runs with a 1.5 min pause in between, allowing subjects to maintain a high level of attention and lessen strain on the eyes. Two high faces and two low faces in each of the three runs were paired with their original temperatures, to prevent extinction. These “booster trials” were not included in the statistical analysis. In experiment 1, the exposure time of the faces during the test sequence was 100 ms. In experiment 2, the exposure time of the faces during the test sequence was 12 ms (one refresh cycle) and then a mask was exposed for 84 ms (seven refresh cycles). The mask consisted of an abstract image that had the same asynchrony (SOA) was 15 s (range 13–17 s). 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The mask consisted of an abstract image that had the same visual properties as the faces, but it was not representing anything more than a number of small squares put together. The same mask was used for all faces in experiment 2. During the calibration, conditioning, and test sequence, the experimental leader was placed in a chair in the back of the room, facing the subject. The experimental leader repeated each verbal pain rating in a monotonous and control voice before recording the ratings. If the subject experienced an incongruity, s (he) was instructed to make a quick correction, e.g., say “No, not fifty, fifty-TEEN.” Only in rare cases (<1%), such corrections were required. Moreover, the placement of the experimental leader allowed for constant monitoring to make sure that subjects’ were truly facing the screen and not looking in other directions.

After the last test sequence of experiment 1, subjects were asked whether the exposure time of the pictures allowed them to see each picture properly. All subjects in experiment 1 reported that they could clearly see the content of the mask and discriminate between the different faces. To verify that the stimuli in experiment 2 would be truly nonrecognizable, we first conducted a methodological pilot study in seven healthy individuals who were exposed to the visual paradigm of experiment 2 and then asked to perform a face recognition test. The instruction was: “You are about to see some pictures on the screen again and I would like you to answer if you have seen this face before during the experiment. You can only say “yes” or “no.” The pictures will be exposed to you very quickly so you might not be able to tell if you saw it before or not. In any case, you have to guess “yes” or “no” for each exposure.” The recognition test included 12 exposures of the previously used faces and 12 exposures of new faces and participants were asked to indicate whether the face had been exposed before, or not. Mean accuracy of identification was 53% (± 10), P = 0.466, confirming that the stimuli were indeed nonrecognizable and thus the parameters were used in experiment 2. Immediately after the test sequence of experiment 2, subjects were asked whether they could recognize the images properly. All subjects in experiment 2 answered that they could not consciously discriminate between the masked faces. To verify this, they were also presented with the face recognition test that was used in the experiment. The accuracy of the recognition test, performed at the end of experiment 2, was analyzed using a repeated measures ANOVA. Results validate that there was no significant difference in recognition for any of the 12 faces used in the face recognition test, for any of the two exposures of each face [six new faces, six old faces, with two exposures each = 24 exposures in total. First exposure, main effect for face type F(1, 18) = 0.073, P = 0.790, nonsignificant; second exposure: main effect for face type F(1, 18) = 0.397, P = 0.538, nonsignificant.] The mean accuracy in the recognition test after experiment 2 was 59.9% (± 10), P = 0.003, one-sample t test. Four individuals had a recognition rate >70%, which contributed to a high mean recognition accuracy. However, the main result of experiment 2 was still significant, even if the subjects with the highest recognition accuracy were removed from the statistical analyses: recognition accuracy 52% (±8), P = 0.392, one-sample t test. Significant main effect for face type in a repeated measures ANOVA was: F(2, 22) = 7.25, P = 0.004, and significant pairwise comparisons between the highflow (P < 0.009), high/low control (P < 0.047) and low/low control (P < 0.019).

ACKNOWLEDGMENTS. We thank Dr. Jonathan Barrebi who provided valuable technical expertise for this study and Prof. Arne Ohman and Dr. Predrag Petrovic for providing valuable theoretical support for this manuscript. The work is supported by funding from the Swedish Society for Medical Research and the Swedish Council for Working Life and Social Research (to K.B.J.) and Grants R21AT004497 (National Center for Complementary and Alternative Medicine, NCCAM), R03AT218317 (National Institute on Drug Abuse), and R01AT006364 (NCCAM) (to J.K.); and R21AT004497 (National Center for Complementary and Alternative Medicine, NCCAM), R03AT218317 (National Institute on Drug Abuse), and R01AT006364 (NCCAM) (to J.K.); K24AT004095 (NCCAM) (to T.J.K.); and R01AT005280 (NCCAM) (to R.L.G.).


