Predictive attenuation of touch and tactile gating are distinct perceptual phenomena

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In recent decades, research on somatosensory perception has led to two important observations. First, self-generated touches that are predicted by voluntary movements become attenuated compared to externally generated touches of the same intensity (attenuation). Second, externally generated touches feel weaker and are more difficult to detect during movement compared to rest (gating). Researchers today often consider gating and attenuation to be the same suppression process; however, this assumption is unwarranted because, despite more than forty years of research, no study has combined them in a single paradigm. We quantified how people perceive self-generated and externally generated touches during movement and rest. We demonstrate that whereas voluntary movement gates the precision of both self-generated and externally generated touch, the amplitude of self-generated touch is selectively attenuated compared to externally generated touch. We further show that attenuation and gating neither interact nor correlate, and we conclude that they represent distinct perceptual phenomena.

Let’s imagine you are at your doctor for a medical examination. Upon her request, you apply pressure with your index finger on your leg to indicate where exactly you feel the pain. The pressure you feel on your leg and the tip of your finger is the feedback from your voluntary finger movement, and it is called somatosensory reafference. To reproduce and confirm your sensations, imagine that the doctor now applies pressure with her index finger at the same spot on your leg. This pressure is generated by the doctor and not from you, and it is called somatosensory exafference. Now imagine that the doctor asks you to first keep your leg relaxed and then flex and extend it while you or her continuously apply pressure on your leg. You can therefore experience your reafferent or her exafferent touches on your leg, while it is moving or resting. Distinguishing between these four conditions is fundamental for your sensorimotor control; your nervous system must know both the source of the touch and the state of your limb to make appropriate use of the sensory feedback. A cutaneous mechanoreceptor in your peripheral nervous system though, cannot distinguish whether a touch is reafferent or exafferent, and thus this distinction must be made centrally, where tactile signals from the skin, sensory information from muscles and joints, and information from motor commands are available. How, then, does the central nervous system classify somatosensory signals during movement?

Several experimental studies in humans have shown that the brain attenuates somatosensory reafference compared to exafference. In behavioral research, this refers to participants perceiving self-generated strokes, forces, or taps as weaker than external equivalents of the same intensity \textsuperscript{1–10}. This somatosensory attenuation has been related to reduced activity in the secondary somatosensory cortex \textsuperscript{11–13} and the cerebellum \textsuperscript{11,13,14} and an increased connectivity between the two areas \textsuperscript{13,15} during self-generated compared to externally generated touches. Somatosensory attenuation was found in 98\% or 315 out of 322 people across a wide age range \textsuperscript{16}, and it is considered one of the reasons why we cannot tickle ourselves \textsuperscript{17–19}. 
Sensory attenuation is not exclusive to humans; similar strategies are used in other species across the animal kingdom (for reviews, see 20–24). For example, during self-chirping, the cricket’s central auditory processing is inhibited (both presynaptically and postsynaptically) in phase with the insect’s chirps to prevent desensitization of its auditory system while maintaining sensitivity to external sounds 25,26. In mice, auditory cortical responses in response to self-generated sounds are attenuated, and this attenuation is present only for the tone frequencies the animal has associated with its locomotion and absent when the same sounds are externally produced 27. A weakly electric fish 28 is able to respond exclusively to externally generated electrical discharges by attenuating its predicted electrosensory reafference 21,29. In primates, activity in the vestibular nucleus in response to vestibular reafference is attenuated during active head movements compared to passive head movements, allowing the animal to maintain its head and body posture and activate the vestibular-related reflexes when appropriate 20,21,30–32.

At the same time, another branch of experimental research has shown that somatosensory sensitivity in response to externally generated stimuli is *gated* during and before a voluntary movement. In human psychophysical research, this phenomenon of *movement-related tactile gating or tactile suppression* manifests as an elevation in the detection threshold 33–42 and a decrease in the subjective intensity of externally generated stimuli 35,37,43 when the stimulated body part moves compared to when it is at rest. Several electrophysiological studies have shown this gating as an inhibition of somatosensory evoked potentials during active movement compared to rest, at subcortical and cortical sites along the somatosensory pathway 43–48. Similar to somatosensory attenuation, tactile gating is a biologically preserved mechanism that is seen across different species 49. For example, responses recorded at the cat’s medial lemniscus evoked by nerve stimulation are suppressed prior to and during limb movements 50. Similarly, the transmission of cutaneous afferent signals to the primary somatosensory cortex is suppressed in rats during movement compared to rest 51. In monkeys, the gating of cutaneous afferent input during active movement has been observed both in the primary somatosensory cortex 52–54 and the spinal cord 54,55.

Somatosensory attenuation and tactile gating share two important conceptual similarities. First, they both refer to a modulation —either in terms of magnitude or precision— in the perception of cutaneous stimuli during movement. Second, they have been assigned the same functional role 39: to reduce the flow of afferent information that can be predicted from the motor command, so that detection of external input that can be biologically important, such as touches caused by predators 17,24,56, or input that is task-relevant for the upcoming or ongoing movement 45,47,57 is facilitated.

Importantly, however, the two phenomena present one striking difference. Somatosensory attenuation relates to somatosensory reafference, that is, touches caused by our voluntary movement. In contrast, gating relates to somatosensory efference, that is, external touches occurring during our voluntary movement. Nevertheless, somatosensory research, including very recent studies and review articles, e.g., 41,42,58–64, often treats the two phenomena as a single generalized suppression strategy of the brain and uses the terms attenuation and gating interchangeably. If the two phenomena are indeed different, this false equivalence is detrimental for understanding human sensorimotor control. First, it cannot advance our understanding at the computational level because researchers try to explain gating (e.g., 41) and attenuation (e.g., 65) using the computational processes proposed for attenuation and gating, respectively. Similarly, at the neurobiological level, researchers intermix neural correlates of gating and attenuation (e.g., 62,63) because they assume that these measure the same single
phenomenon. Second, this confusion becomes particularly disadvantageous in clinical studies using gating and attenuation when interpreting findings of sensorimotor deficits in patients with schizophrenia 66–68, functional movement disorders 69 or Parkinson’s disease 45,70,71.

Are these phenomena the same, or do they represent two fundamentally distinct processes? Does the brain treat all sensory stimuli similarly during movement, independent of whether they are reafferent or exafferent? To the best of our knowledge, there has been no previous attempt to simultaneously test attenuation and gating with the same stimulus and psychophysics task. Here, in a single experimental design, we investigated the perception of touches applied on the left hand while manipulating whether the left arm was in movement or resting (left limb state). We additionally manipulated whether the touches were reafferent, generated by the right hand, or exafferent, generated by an external source (origin of touch). We reasoned that if the two phenomena are the same, then they should influence somatosensory perception in similar ways. Our results demonstrate that this is not the case: voluntary movement reduces the somatosensory precision of both sensory reafference and exafferece (gating), while the perceived amplitude of sensory reafference is selectively attenuated compared to exafference (attenuation). Notably, the two phenomena do not correlate with each other or interact when present simultaneously. Thus, collectively, our results show that gating and attenuation are two separate processes that can be experimentally dissociated.

Results
Participants rested their left hands palm up with their index fingers placed on a molded support, and their right hands were palm down on top of a set of sponges (Fig. 1a-d). In all conditions, they performed a force-discrimination task 3,4,10,67: in each trial, a motor delivered two taps (the test tap and the comparison tap) on the pulp of their left index finger, and they were asked to verbally indicate which tap felt stronger (Fig. 1e-h). While the test tap had a fixed intensity (2 N), the comparison tap randomly changed its intensity on every trial (1, 1.5, 1.75, 2, 2.25, 2.5, or 3 N). An auditory ‘go’ signal indicated the trial’s onset and the onset of the response period.

In all conditions, our experimental manipulation exclusively concerned the test tap; the comparison tap was always externally triggered and delivered on the relaxed left arm, serving, therefore, as a reference stimulus. In a factorial design, we controlled for whether the left arm moved (Fig. 1c, d) or remained relaxed (Fig. 1a, b) during the test tap, and whether the test tap was produced by the right hand (self-generated) (Fig. 1b, d) or not (externally generated) (Fig. 1a, c). This design resulted in four experimental conditions, the order of which was fully counterbalanced (Fig. 1a-d).

In the baseline condition (Fig. 1a), participants did not move their limbs but passively received the test and the comparison taps on the left index. This control condition assesses the participants’ somatosensory perception in the absence of any movement 2,4,10. In the attenuation condition (Fig. 1b), participants actively tapped with their right index finger a force sensor placed on top of their left index finger. The tap of their right index finger on the force sensor triggered the test tap on their left index finger. This classic condition assesses the perception of a self-generated tap on a passive limb 2,4,10. In the gating condition (Fig. 1c), participants were asked to continuously move their left arm forward, sliding the experimental setup with the motor and the force sensors between a start and an end position (distance 25 cm) at a comfortable velocity of approximately 20 cm/sec (Fig. 1i-j, Supplementary Fig. S1, S2). During this movement, participants received the test tap on their left index finger. This is a gating condition because it assesses the perception of an externally generated tap on a moving
limb. Finally, in the attenuation&gating condition (Fig. 1d), the participants performed the same movement with their left arm, but they were additionally asked to actively tap with their right index finger on the force sensor that triggered the test tap on their left index finger during the movement. The force sensor was attached to the experimental setup and moved together with the left hand. This condition combines the gating and attenuation phenomena since it assesses the perception of a self-generated tap on a moving limb.

Fig. 1. Experimental conditions. Two factors were manipulated in the experiment, resulting in four experimental conditions: whether the left arm was in rest (a, b) or in movement while receiving the test tap (c, d) (state of the left limb) and whether the test tap was externally triggered by the motor (a, c) or self-triggered by the participants’ right hand (b, d) (origin of touch). In all four conditions (e-h), the participants received the two taps (test and comparison tap) on the pulp of their left index fingers from the motor (black stars), and they had to verbally indicate which was stronger: the first or the second tap. In the attenuation and attenuation&gating conditions, the participants self-triggered the test tap on their left index finger by moving their right arm to tap a sensor with their right index finger (blue stars). (i) In the gating and attenuation&gating conditions (c and d), the participants extended their left arm from a start position to an end position, sliding the experimental setup along two drawer runners. During this movement, participants experienced the test tap. A motion sensor recorded the position of the platform in time. (f) Example position traces recorded by the motion sensor for the movements of one participant during the attenuation&gating condition. (g) Responses of a representative participant and fitted logistic
models for the four experimental conditions. Data points have been horizontally jittered to avoid overlapping.

The participant’s responses in each condition were fitted with a generalized linear model (Fig. 1k, Supplementary Fig. S3), and two parameters of interest were extracted: the point of subjective equality (PSE), which represents the intensity at which the test tap felt as strong as the comparison tap \( p = 0.5 \), and the just noticeable difference (JND), which reflects the participants’ sensitivity in force discrimination. We hypothesized that the two phenomena are different and, thus, that they would affect the PSE and JND differently. Specifically, we expected a decrease in the perceived magnitude (lower PSE) for the conditions with sensory reafference (attenuation and attenuation&gating conditions), with no effects on the sensory precision of the participants (JND) when they received the reafference on a still limb (attenuation condition). In contrast, we predicted a decrease in the sensory precision (higher JND) of both sensory reafference and exafference for the conditions where the limb that receives the touches moves (gating, attenuation&gating) and small, if any, effects in the PSE. In other words, we expected that the attenuation phenomenon affects the PSE and not the JND, and the gating phenomenon mainly affects the JND. Our hypotheses were supported by the data (Fig. 2a-g).

**Fig. 2.** Experimental results. (a, b) Bar graphs show the PSES and JNDs (mean ± SEM) for each condition. A lower PSE value indicates a lower perceived magnitude, while a higher JND value indicates a lower somatosensory sensory precision. Reafferent touches produced by the movement of the right arm (attenuation and attenuation&gating conditions) were associated with a significant drop in the PSES, while the movement of the left arm that receives the touches (gating and attenuation&gating) produces a significant increase in the JNDs. (c) Group psychometric functions for each condition generated using the mean PSE and the mean JND across participants. The leftward shift of the curves...
for the attenuation and attenuation\&gating conditions illustrates that somatosensory reaффerence is perceived as weaker than aффerence. The flattening of the curves for the gating and attenuation\&gating conditions illustrates the worsening in somatosensory precision of both aффerence and reaффerence on a moving limb. (d, e) Line plots illustrate the decreases in PSEs when experiencing reaффerent touches compared to aффerent touches, when the left arm is still (d) and when the left arm moves (e). (f, g) Line plots illustrate the increases in JNDs when receiving the touches on a moving limb compared to rest, when the touches are aффerent (f) and when the touches are reaффerent (g).

We performed a repeated-measures ANOVA on the PSEs with the origin of the touch (reaффerence vs aффerence) and the state of the left limb (movement vs rest) as factors. This revealed a significant main effect of the origin of the touch \((F(1, 23) = 36.10, p < 0.001, \eta^2_p = 0.611)\), a significant main effect of the left limb state \((F(1, 23) = 13.91, p = 0.001, \eta^2_p = 0.377)\) and a nonsignificant interaction \((F(1, 23) = 0.26, p = 0.615, \eta^2_p = 0.011)\) (Fig. 2a). A Bayesian repeated-measures ANOVA further concluded against the interaction term, by supporting the model without the interaction term compared to the full factorial \((BF_{M1} / BF_{M2} = 3.631)\). The attenuation condition produced a significant drop in the PSE compared to the baseline condition \((n = 24, t(23) = -5.908, p < 0.001, \text{Cohen’s } d = -1.206, CI^{95} = [-0.332, -0.160], BF_{01} < 0.0003)\) (Fig. 2d). This replicates previous attenuation findings indicating that a self-generated tap on a passive limb feels weaker than a tap of the same intensity but of an external origin \(^{2,4-7,10,13,16}\). Similarly, the PSE in the attenuation\&gating condition was significantly lower than that in the gating condition \((n = 24, t(23) = -5.032, p < 0.001, \text{Cohen’s } d = -1.027, CI^{95} = [-0.377, -0.157], BF_{01} = 0.002)\) (Fig. 2e), extending the previous conclusion to when the receiving limb is moving. Together, these two contrasts show that reaффerent (self-generated) touches feel weaker than aффerent touches, both when the receiving hand is in movement or rest.

The gating and attenuation\&gating conditions also produced a significant drop in the PSE compared to the baseline condition \((n = 24, t(23) = -2.409, p = 0.024, \text{Cohen’s } d = -0.492, CI^{95} = [-0.141, -0.011])\) and the attenuation condition \((n = 24, V = 55, p = 0.005, rrb = -0.633, CI^{95} = [-0.161, -0.022])\), respectively. However, these decreases were quite modest (\(\approx 30\%\) of the weakening produced by the attenuation condition) and supported only by anecdotal evidence from Bayesian statistics \((BF_{01} = 0.433 \text{ and } BF_{01} = 0.738, \text{respectively})\). Together, these contrasts suggest that aффerent touches can feel slightly weaker on a moving limb than on a passive limb, in agreement with previous findings for tactile gating \(^{35,37,43}\). Nevertheless, when compared with tactile reaффerence, the perceived magnitude of tactile reaффerence is not strongly decreased.

When testing for the effects of the conditions on the somatosensory precision of the participants (JND), there was a significant main effect of the state of the left limb \((F(1, 23) = 17.1, p < 0.001, \eta^2_p = 0.426)\), but there was neither a significant main effect of the origin of touch \((F(1, 23) = 0.52, p = 0.478, \eta^2_p = 0.022)\) nor a significant interaction \((F(1, 23) = 0.06, p = 0.809, \eta^2_p = 0.003)\) (Fig. 2b). As with PSEs, the absence of interaction for the JNDs was further supported with a Bayesian repeated-measures ANOVA that provided evidence against the interaction term \((BF_{M1} / BF_{M2} = 3.522)\). The attenuation condition did not produce any change in the JND with respect to the baseline condition \((n = 24, t(23) = 0.331, p = 0.744, \text{Cohen’s } d = 0.068, CI^{95} = [-0.034, 0.047])\), and this was substantially confirmed with a Bayesian analysis \((BF_{01} = 4.432)\). In alignment, no significant differences in the JND were detected between the gating and the attenuation&gating conditions \((n = 24, t(23) = 0.72, p = 0.481, \text{Cohen’s } d = 0.146, CI^{95} = [-0.024, 0.05])\), and this was again confirmed with a Bayesian analysis \((BF_{01} = 3.691)\). Together, these two contrasts demonstrate that receiving sensory reaффerence per se is not accompanied by a worsening of sensory precision on the receiving limb.
In contrast, moving the limb while receiving an external touch (gating condition) produces a significant increase in the JND compared to the baseline condition \((n = 24, t(23) = 3.134, p = 0.005, \text{Cohen's } d = 0.640, CI^{95} = [0.019, 0.093], BF_{01} = 0.108)\) (Fig. 2f). This was further confirmed by a significant increase in JND between the attenuation and the attenuation&gating conditions \((n = 24, t(23) = 2.984, p = 0.007, \text{Cohen's } d = 0.609, CI^{95} = [0.019, 0.106], BF_{01} = 0.146)\) (Fig. 2g). Together, these two contrasts indicate that voluntary movement per se deteriorates the precision with which reafferent and exafferent stimuli are perceived on the moving limb.

Together, our results indicate that predicting the sensory consequences of a voluntary movement produces a decrease in the perceived magnitude of sensory reafference (PSE) without a concomitant worsening in somatosensory precision (JND). In contrast, a voluntary movement leads to a decrease in somatosensory precision (JND) for both sensory reafference and exafference. These effects are observed in the group psychometric fits (Fig. 2c). To further illustrate that somatosensory attenuation affects the amplitude (PSE), while tactile gating affects the precision (JND) and not vice versa, we tested whether we could predict better the participants’ performance in the attenuation&gating condition when using the PSE from the attenuation condition and the JND from the gating condition than the PSE from the gating condition and the JND from the attenuation condition. Indeed, the first model was significantly better: \(n = 24, V = 39, p < 0.001, rrb = -0.74, CI^{95} = [-68.980, -12.736], BF_{01} = 0.443\) (Supplementary Fig. S4, Fig. 3a-b).

In the abovementioned ANOVAs, there were no significant interactions between the two factors (the origin of touch and the state of the limb), neither for the PSEs nor for the JNDs, according to both frequentist and Bayesian analyses. This indicates that the effect of the left limb state was not influenced by the effect of the origin of touch, neither for the PSE nor for the JND. Instead, the two effects just added up when simultaneously present. To further test whether the effects produced by each phenomenon relate to each other at all, we performed a correlation analysis across the PSEs and JNDs. No significant correlations were detected between any of the PSEs and any of the JNDs (all \(p\)-values > 0.225, \(BF_{01} = [1.971, 3.950]\)). The only significant correlation found within PSEs was between the PSE in the attenuation condition and the PSE in the attenuation&gating condition \((t(22) = 4.89, r = 0.722, p < 0.001, CI^{95} = [0.449, 0.871], BF_{01} = 0.002)\) (Fig. 3c). That is, the weaker the participants perceived the magnitude of their self-generated touch during rest, the weaker the magnitude of their self-generated touch during movement felt. Given that the PSEs significantly dropped in these two conditions and that these decreases correlated with each other, this result provides further support that their common experimental denominator, i.e., the reafferent nature of the touch, was responsible for the drop in the PSE and thus for the attenuation phenomenon. In contrast, the JND in the gating condition was significantly correlated only with the attenuation&gating condition \((t(22) = 3.47, r = 0.595, p = 0.008, CI^{95} = [0.252, 0.805], BF_{01} = 0.047)\) (Fig. 3d). This specific correlation means that the worse the somatosensory precision of an external touch when participants moved their receiving hand, the worse the sensory precision for a self-generated touch during the same movement of the receiving hand. Given that the JNDs significantly increased only in these two conditions and that these increases correlated with each other, this result provides evidence that their common experimental denominator, i.e., the left limb's movement that receives the touches, was responsible for the increase in the JND.
Fig. 3. Model predictions and scatterplots for PSEs and JNDs. (a) Average participants’ responses in the attenuation&gating condition (yellow) and average predicted responses using the parameters from the attenuation and gating conditions (blue). The responses depicted in dark blue used the PSE of the attenuation condition and the JND from the gating condition, while the responses depicted in light blue used the PSE of the gating condition and the JND of the attenuation condition. Error bars and ribbons represent 95% confidence intervals. (b) For each participant, we estimated the Bayesian Information Criterion (BIC) of the fitted logistic model in the attenuation&gating condition and the two models with fixed parameters from the attenuation and gating conditions. The model using the PSE of the attenuation condition and the JND of the gating condition was a significantly better model than the one using the PSE of the gating condition and the JND from the attenuation condition. The upper right panel represents the same data after excluding one participant corresponding to the outlier observation, for illustration purposes. The exclusion of the outlier did not change the statistical results. (c) The participants’ PSEs in the attenuation condition were significantly correlated with those in the attenuation&gating condition. (d) The participants’ JNDs in the gating condition were significantly correlated with those in the attenuation&gating condition. There were no significant correlations between PSEs and JNDs.

A common finding in tactile gating studies is that the gating effects are stronger with higher movement velocities \(33,36,40,41\); that is, the faster the limb movements are, the worst the
perception of the moving limb. Therefore, one could hypothesize that any differences observed
between the gating and the attenuation\&gating conditions might be due to differences in the
velocity of the participants’ movements. Since there were no significant JND differences in
these two conditions (Fig. 2), which was further supported by a Bayesian analysis, this concern
can be ruled out. However, one can argue that the PSE in the attenuation\&gating condition was
lower than the PSE in the attenuation condition because the participants moved faster in the
attenuation\&gating condition and not because of the reafferent nature of the touch. This
concern can also be ruled out since, rather than faster, we observed that the participants moved
slightly slower in the attenuation\&gating condition (20.3 ± 0.003 cm/s) than in the gating
condition (23.3 ± 0.003 cm/s); (peak trial velocity; t(23) = -4.062, p <0.001, Cohen’s d = -
0.829, CI\(^95\) = [-0.001, -0.004], BF\(_{01} = 0.015\)). Although the total distances the participants ran
with their left arm were comparable, participants in the attenuation\&gating condition moved
slower because they had to coordinate both their arms to tap the sensor that the left arm moves
with the right hand (Fig. 1d). This difference was further confirmed when looking at the peak
velocities during the test tap between the two conditions (gating: 22.2 ± 0.003 cm/s;
attenuation\&gating: 19 ± 0.003 cm/s) (Supplementary Fig. S5). Therefore, these kinematic
differences cannot explain the significant decrease in the PSEs in the attenuation\&gating
condition compared to the gating condition, but they actually underscore the importance of
reafference in somatosensory attenuation.

Discussion

The present study contrasted the conditions of attenuation and gating in a single experimental
paradigm to investigate the relationship between the two phenomena. To this end, we
independently manipulated the origin of the touch (reafference vs exafference) and the state of
the receiving limb (movement vs rest), taking on all four possible combinations of these levels.
We replicated the classic phenomenon of predictive attenuation of touch 1,2,6,10,3,34 by showing
that somatosensory reafference feels weaker than somatosensory exafference. Importantly,
however, this decrease in the perceived amplitude (PSE) was not accompanied by a
concomitant worsening in somatosensory precision (JND). Participants had the same
discrimination capacity for both reaferent and exafferent touches applied on their passive limb,
a conclusion supported by Bayesian statistics. Nevertheless, when the limb that receives the
touches moved, this voluntary movement per se led to a decrease in somatosensory precision
(JND) for both reafferent and exafferent touches, replicating the classic tactile gating
phenomenon 33,36,39,42. The two effects did not correlate and did not interact but summed up
when present together.

The main conclusion of the present study is that the predictive attenuation of touch and tactile
gating are two distinct perceptual phenomena. Our findings can help conciliate several previous
observations on gating and attenuation that have been studied in isolation during recent decades.
First, attenuation is observed not only on the active limb 8 but also on a passive limb (the
contralateral hand in the present study), as long as the contact between the body parts is
predicted by the voluntary movement 2,7,9,10,13,16. In contrast, there is abundant evidence that
external touches applied to the limb contralateral to the one that moves are not gated 34,43,45,72,73.
Second, a touch that results from a passive movement 10 or touches that are simultaneously
presented in both hands (double touch) 67 are not attenuated. In contrast, gating effects have
been repeatedly demonstrated for passive movements, both electrophysiologically 45 and
behaviorally 34,38,39. Third, self-generated tactile signals are attenuated as long as they are
presented at the timing predicted by the action 1,2,4; even a 100 ms delay between the movement
and its tactile feedback greatly reduces the attenuation of the latter. In contrast, externally generated stimuli are gated when presented at any (unpredicted) timing during the movement, and importantly, even at hundreds of milliseconds before the movement onset. Fourth, whereas gating shows no specificity for the type of motor activity and manifests both during isotonic and isometric contractions, attenuation is motor command-specific; a consistent but arbitrary and unnatural mapping between the motor command and the touch, for example, moving a joystick with one hand to produce touch on the other, does not produce attenuation. Together, when seen in light of our findings, these results are not contradictory; instead, they refer to different perceptual phenomena.

Motor control relies on integrating afferent sensory information with efferent motor signals. Distinguishing between gating and attenuation is fundamentally important for motor control theories because this can indicate a different integration or weighting mechanism of the motor and sensory information depending on the context. One well-established computational framework inspired by engineering approaches poses that the brain produces the motor command through an inverse model or controller. A copy of the motor command — termed ‘efference copy’ — is used by a forward model to predict the expected sensory feedback of the movement, which is then combined with the actual sensory input to estimate the state of the body. With respect to the attenuation of sensory reafference, it has been proposed that the prediction signal of the forward model is used to ‘cancel’ the sensory reafference. In other words, central motor processes play a major role in somatosensory attenuation compared to the actual sensory feedback. Support for this comes from studies showing that conditions that present highly predictable touches but in the absence of movement do not yield attenuation (see also for similar conclusions). The dependence of attenuation on action prediction was further shown when participants attenuated touches applied on one hand that were predicted by their other hand’s movement, even when the two hands unexpectedly failed to make contact. In further agreement, neuroimaging studies on somatosensory attenuation consistently report activation of the cerebellum, a structure that is associated with motor prediction.

In contrast, this computational account that relies on action prediction, efference copy, and internal forward models is inapplicable for tactile gating, since gated touches can be of exafferent nature and occur at any (unpredictable) time during movement, even before movement onset. In other words, there is no information that the brain can use to predict the exafferent touches with the forward model because there is no causal relationship between the motor command and sensory input. This observation agrees with the proposal that peripheral afferent signals from muscle spindles and joint afferents play the major role in gating, and the fact that gating effects have also been observed during passive movement, without significant differences from active movements. Then, if efference copy is not the basis for gating, how can the gating effects be computationally explained?

The alternative computational framework of active inference has been proposed to explain somatosensory attenuation; however, based on an assumed equivalence between attenuation and gating. The active inference approach refutes the necessity of an efference copy, and emphasizes the importance of a generative model and reflex arcs in the place of forward and inverse models, and controllers. According to the active inference account, the brain predicts the sensory input that would be expected from a specific action, and the body moves in order to fulfill these sensory predictions. Motor commands are thus conceptually replaced by proprioceptive predictions, and action occurs as a way to minimize the proprioceptive prediction errors when the movement has not been yet executed. A major role is assigned to...
the precision (i.e., reliability) that weights these sensory prediction errors depending on the context, and can be manipulated through attention allocation. Within this computational architecture, attenuation of somatosensory reafference is seen as a reduction in the precision of somatosensory evidence during movement in order to allow the expression of proprioceptive predictions that trigger the movement 59. In other words, the agent attends away from somatosensory input in order to execute the movement. However, this proposal does not address the attenuation of sensory reafference with respect to exafference, since the agent should theoretically attend away from all somatosensory input, independently of their source. Moreover, the proposal cannot explain why increasing the precision of the proprioceptive prediction errors on the hand that is to move would reduce the precision of somatosensory prediction errors on the contralateral limb that is not meant to move and thus, there should not exist any proprioceptive predictions about it. In contrast, the active inference account can sufficiently explain the tactile gating effect; i.e., the reduction in the precision of somatosensory input on the moving limb during the movement.

A recent study 64 suggested that action prediction does not produce somatosensory attenuation but enhancement and argued that the attenuation observed in all earlier experiments is actually due to nonpredictive gating processes. This claim cannot be supported based on several observations and arguments. First, the present results showed that the predictive attenuation of touch and gating are different phenomena; if sensory attenuation was the same as tactile gating, then all stimuli applied on the moving limb would be attenuated. In contrast, we show that sensory reafference is selectively attenuated compared to sensory exafference, both in a passive and a moving limb. Similar results were shown in 79, where participants attenuated only their reafferent touches and not exafferent touches presented at the same limb simultaneously.

Second, Thomas and colleagues 64 provided participants with an arbitrary mapping between the movement of one hand and sensory feedback on the other hand. Using a very similar setup, a study from 2006 3 found neither attenuation, which is consistent with the predictive account of attenuation, nor enhancement effects, which is inconsistent with those from Thomas and colleagues 64. Third, it is essential to keep in mind that attenuation has also been shown on an anesthetized limb 8 and for imagined movements 9, further emphasizing the contribution of motor prediction in relation to peripheral feedback for somatosensory attenuation. Finally, it has been shown that when reaching with one arm towards the other, the perceived intensity of an external tactile stimulus applied to the passive target limb is enhanced 92. Although we believe that future experimentation is needed to understand and replicate the reported enhancement effects 64, we speculate that, as in 92, increased spatial attention or other kinematic differences play a significant role in the somatosensory enhancement reported by Thomas and colleagues 64.

We conclude that the human brain uses two different basic processes to suppress reafferent and exafferent information during movement and rest. This separation of attenuation and gating can explain why although we cannot tickle ourselves because we attenuate our self-tickles, we do tense our muscles when being tickled by others to decrease our sensitivity to the external tickles.

Materials and Methods

Participants

After providing written informed consent, twenty-four participants (12 women and 12 men, 22 right-handed, 1 ambidextrous, 1 left-handed) aged 21-40 years participated. Handedness was assessed using the Edinburgh Handedness Inventory 93. The sample size was set to twenty-four (24) before data collection commenced based on our previous studies using the same methods.
while ensuring a counterbalanced order of conditions. The Swedish Ethical Review Authority (https://etikprovningsmyndigheten.se/) approved the study (no. 2016/445-31/2, amendment 2019-04536). All participants provided their written informed consent.

General Procedure

Participants sat comfortably on a chair with their arms placed on a table. Their left hands rested palm up, with their index fingers placed on a molded support. The right arms rested palm down on top of a set of sponges. In each trial, a motor (Maxon EC Motor EC 90 flat; Switzerland) delivered two taps (the test tap and the comparison tap) on the pulp of their left index finger through a cylindrical probe (25 mm height) with a flat aluminum surface (20 mm diameter) attached to the motor’s lever. A force sensor (FSG15N1A, Honeywell Inc.; diameter, 5 mm; minimum resolution, 0.01 N; response time, 1 ms; measurement range, 0–15 N) within the probe recorded the forces applied on the left index finger. Following the presentation of the two taps, participants had to verbally indicate which tap felt stronger: the first or the second.

A wooden surface was placed under the motor and the sensors. This surface was placed on top of two commercially available drawer runners (IKEA, https://www.ikea.com/us/en/p/besta-drawer-runner-soft-closing-40348715/). One side of the runners was attached to the table with Velcro material, and the other side was attached to the bottom side of the surface. With this configuration, the surface – with the motor, the sensors and the participants’ hands – could be moved forward and backward.

In all conditions, the comparison tap was delivered on the left index finger with a random delay of 800-1500 ms from the test tap. In the attenuation and attenuation&gating conditions (Fig. 1b, d), the tap of the participants’ right index fingers on the force sensor triggered the test tap on their left index finger with an intrinsic delay of »36 ms.

In the gating and attenuation&gating conditions (Fig. 1c, d), participants were asked to extend their elbow upon the auditory ‘go’ cue. The extension of the elbow moved the platform forward on the table (Fig. 1f). A green tape on the table (Fig. 1i) indicated the start position of the platform, while a red tape indicated its end position. The participants were asked to move the platform from the start position to the end position (distance = 25 cm). During the movement of the left arm, the participants received the test tap on their left index finger. Before the condition started, we emphasized to the subjects that their task was to pay attention to the force that they would receive during the movement, rather than covering exactly the distance between the lines. Moreover, the participants were trained to perform the movement in approximately 1000-1500 ms after the ‘go’ cue and then stop. In the gating condition, the test tap was applied 800 ms after the ‘go’ cue so that its delivery was during the movement. Similarly, in the attenuation&gating condition, the participants triggered the test tap during the movement. The comparison tap was applied 800-1500 ms after the test tap to ensure that the participants had stopped moving. Once the participants gave a response, they returned the platform to the start position.

A motion tracking sensor (6DOF Polhemus Fastrak, USA, weight = 9.1 g, dimensions = 2.29 cm x 2.82 cm x 1.52 cm) was placed on top of the platform to record the motion of the platform due to the movement of the participants’ left arm. The sensor recorded the x, y and z positions at a sampling rate of approximately 120 Hz.

Each condition included 70 trials. The test tap was set to 2 N, while the intensity of the comparison tap was systematically varied among seven different force levels (1, 1.5, 1.75, 2,
Each tap lasted for 100 ms. On every trial, participants verbally indicated which tap on their left index finger felt stronger: the first (test) or the second (comparison). Participants were told not to try to balance their responses (50% first and 50% second), and they were further instructed to make their best guess if the intensity of the two taps felt similar.

In addition, participants were administered white noise through a pair of headphones to preclude any sounds created by the motor to serve as a cue for the task. The loudness of the white noise was adjusted so that participants could clearly hear the auditory cues of the trial. In all conditions, the view of the pulp of the left index finger was occluded. Participants were asked to fixate on a cross placed on a wall 2 m opposite them, but they were allowed to look at the force sensor to guide the movement of the right index finger when needed (Fig. 1b, d). No feedback was provided to the participants about their responses.

**Force discrimination analysis**

In each condition, the participants’ responses were fitted with a generalized linear model using a logit link function (Equation 1):

\[
p = \frac{e^{\beta_0 + \beta_1 x}}{1 + e^{\beta_0 + \beta_1 x}} \quad \text{(Equation 1)}
\]

We extracted two parameters of interest: the PSE, which represents the intensity at which the test tap felt as strong as the comparison tap \((p = 0.5)\) and which quantifies the perceived intensity, and the JND, which reflects the participants’ discrimination capacity. Before fitting the responses, the values of the applied comparison taps were binned to the closest value with respect to their theoretical values (1, 1.5, 1.75, 2, 2.25, 2.5 or 3 N).

**Kinematic analysis.**

Both position and velocity data were smoothed with a moving average filter in MATLAB 2018a. Velocity was calculated as the first derivative of position. To calculate the distance participants ran in every trial of each condition, we calculated the minimum and the maximum position of the platform during the entire trial duration. The peak trial velocity was defined as the peak velocity of the entire trial. The peak tap velocity was defined as the peak velocity during the period that the test tap was applied.

**Rejection of trials**

After data collection, one hundred seventy-three (173) trials out of 6720 (2.57%) were rejected. First, in thirty-four trials (34), the intensity of the test tap (2 N) was not applied accurately (test tap < 1.85 N or test tap > 2.15 N), and in sixteen (16) trials, the responses were missing. Second, we rejected one hundred seventeen (117) trials in total from the gating and attenuation&gating conditions because participants either did not move their left arm (or moved it too slowly) during the test tap (mean velocity < 10 cm/s) or they moved it during the comparison tap (mean velocity > 5 cm/s). The thresholds were based on a previous study showing no gating effects for velocities smaller than 5 cm/s. The analysis was therefore performed with 6547 trials in total.

**Statistical analysis**

We used R ⁹⁴ and JASP ⁹⁵ to analyze our data. The data normality was assessed using the Shapiro-Wilk test. Depending on the data normality, we then performed planned comparisons using either a paired t-test or a Wilcoxon signed-rank test. We report 95% confidence intervals \((CI^{95})\) for each statistical test. Effect sizes are given by the partial eta-squared \((\eta_p^2)\) for the ANOVAs, by the Cohen’s \(d\) for t-tests or by the matched rank biserial correlation \(rrb\) for the Wilcoxon signed-rank tests. In addition, a Bayesian factor analysis using default Cauchy priors
with a scale of 0.707 was carried out for all statistical tests to provide information about the level of support for the null hypothesis compared to the alternative hypothesis ($BF_{01}$) given the data. Finally, correlations were performed using Pearson’s coefficient $r$ given that the data were normally distributed. All tests were two-tailed.

**Corrections for multiple comparisons**

Since our PSE and JND comparisons were planned, we did not apply corrections for multiple comparisons. However, all results remained exactly the same when applying corrections for false discovery rate (FDR) $^9$. In the correlation analyses, we did apply corrections for multiple comparisons (FDR) since, although we expected correlations between the PSEs and between the JNDs, we had no *a priori* hypotheses for correlations between PSEs and JNDs.

**Acknowledgements**

We thank L. Miller, M. Hauser and M. Chancel for their feedback on an earlier version of the manuscript. K.K. was supported by the Swedish Research Council (VR Starting Grant 2019-01909 granted to K.K.). This work was supported by the Swedish Research Council, Torsten Söderbergs Stiftelse, and Göran Gustafsson’s Stiftelse. We thank Kanaka Raghavan for help with the illustrations.

**Conflicts of interest/Competing interests**

The authors declare no competing financial interests.

**Authors’ contributions**

K.K. and H.H.E. conceived and designed the experiment. K.K. collected the data and conducted the statistical analysis. K.K. and H.H.E. wrote the manuscript.

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