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Published in:
Journal of Neurophysiology

DOI (link to publication from Publisher):
[10.1152/jn.00155.2021](https://doi.org/10.1152/jn.00155.2021)

Publication date:
2021

Document Version
Accepted author manuscript, peer reviewed version

[Link to publication from Aalborg University](#)

Citation for published version (APA):
Henrich, M. C., Frahm, K. S., & Andersen, O. K. (2021). Tempo-spatial integration of nociceptive stimuli assessed via the nociceptive withdrawal reflex in healthy humans. *Journal of Neurophysiology*, 126(2), 373-382. <https://doi.org/10.1152/jn.00155.2021>

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Tempo-spatial integration of nociceptive stimuli assessed via the nociceptive withdrawal reflex in healthy humans.

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Running head: Spinal tempo-spatial integration of nociception via the NWR

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- Data acquisition: M.C.H.
- Data analyses and interpretation: M.C.H., K.S.F. and O.K.A.
- Manuscript draft, revision, and approval: M.C.H., K.S.F. and O.K.A.

Keywords: Nociceptive Withdrawal Reflex, Temporal Summation, Spatial Summation, Nociception

27 **ABSTRACT**

28 Spatial information of nociceptive stimuli applied in the skin of healthy humans is integrated in the spinal
29 cord to determine the appropriate withdrawal reflex response. Double-simultaneous stimulus applied in
30 different skin sites are integrated, eliciting a larger reflex response. The temporal characteristics of the stimuli
31 also modulate the reflex e.g. by temporal summation. The primary aim of this study was to investigate how
32 the combined tempo-spatial aspects of two stimuli are integrated in the nociceptive system. This was
33 investigated by delivering single and double simultaneous stimulation, and sequential stimulation with
34 different inter-stimulus intervals (ISIs ranging 30-500 ms.) to the sole of the foot of fifteen healthy subjects.
35 The primary outcome measure was the size of the nociceptive withdrawal reflex (NWR) recorded from the
36 Tibialis Anterior (TA) and Biceps Femoris (BF) muscles. Pain intensity was measured using an NRS scale.
37 Results showed spatial summation in both TA and BF when delivering simultaneous stimulation.
38 Simultaneous stimulation provoked larger reflexes than sequential stimulation in TA, but not in BF. Larger
39 ISIs elicited significantly larger reflexes in TA, while the opposite pattern occurred in BF. This differential
40 modulation between proximal and distal muscles suggests the presence of spinal circuits eliciting a functional
41 reflex response based on the specific tempo-spatial characteristics of a noxious stimulus. No modulation was
42 observed in pain intensity ratings across ISIs. Absence of modulation in the pain intensity ratings argues for
43 an integrative mechanism located within the spinal cord governed by a need for efficient withdrawal from a
44 potentially harmful stimulus.

45

46 **NEW & NOTEWORTHY**

47 Tempo-spatial integration of electrical noxious stimuli was studied using the nociceptive withdrawal reflex
48 and a perceived intensity. Tibialis Anterior and Biceps Femoris muscles were differentially modulated by the
49 temporal characteristics of the stimuli and stimulated sites. These findings suggest that spinal neurons are
50 playing an important role in the tempo-spatial integration of nociceptive information, leading to a reflex

51 response which is distributed across multiple spinal cord segments and governed by an efficient defensive
52 withdrawal strategy.

53

54 **INTRODUCTION**

55 The NWR was originally described as an elicited motor response characterized by a stereotyped flexion of
56 one or several joints aiming at removing the limb from a potentially tissue damaging stimulus (1). The
57 withdrawal was interpreted as the activation of synergistic muscles to produce flexion of the exposed limb,
58 including the inhibition of antagonist muscles and a compensating opposite process in the contralateral limb
59 to preserve balance (1). More recently, evidence has suggested a functional reflex organization in which the
60 NWR involves flexion and extension of relevant joints, by recruiting multiple groups of muscles, depending
61 on the site being stimulated (2–6).

62 Spatial integration of the nociceptive stimuli into a reflex response might involve processing in supraspinal
63 centers. Primary afferents that depolarize due to a nociceptive stimulus will first synapse in the dorsal horn
64 of the spinal cord. Further processing likely happens through a complex network of inhibitory and excitatory
65 interneurons within the dorsal spinal cord(7). Then, ascending information is projected via the anterolateral
66 system to supraspinal structures (brainstem, thalamus and cortex) eventually producing the perception of
67 pain(8). In parallel, information transmitted to the ventral horn of the spinal cord can lead to the elicitation
68 of the protective NWR. Neurons descending from superior structures (primarily the rostroventral medulla in
69 the brainstem) synapsing in the dorsal horn of the spinal cord have been reported (see 9 for a review) and
70 form the basis of the supraspinal descending excitatory and inhibitory modulation maintaining homeostasis
71 of the spinal nociceptive processing.

72 Previous evidence from animal studies suggested that deep dorsal horn neurons encode the organization of
73 the NWR in modules that allow the coordination of muscle recruitment to generate optimal withdrawal (10–
74 13). Human studies have also supported the presence of this spinal modular organization describing the NWR

75 patterns elicited by stimulation in several sites of the sole of the foot (2). In this context, the term Reflex
76 Receptive Field (RRF) was implemented to define the area of the skin from which a suprathreshold stimulus
77 triggers the recruitment of a certain muscle or group of synergistic muscles (2, 5). The RRFs described in the
78 sole of the foot of healthy humans suggest that the NWR pathway integrate spatial information of the
79 stimulus to elaborate an optimal defensive response. It remains unknown, to what degree this spatial
80 integration is altered when introducing a secondary stimulus shortly after the first. It is known that both the
81 spatial characteristics of the stimulus modulate the NWR, but so does the temporal characteristics. By
82 delivering repetitive electrical stimulation to the same site on the sole of the foot, several studies have shown
83 that temporal summation can modulate the NWR. The magnitude of the NWR is significantly increased with
84 repetitive input (14–20), showing a reflex behavior that resembles the wind-up phenomenon observed in
85 single neuronal recordings in animal studies (21–23). To what extent the temporal summation can be
86 translated into tempo-spatial integration has not been studied. By clarifying this, it may shed light on the
87 spinal tempo-spatial integration of nociceptive input.

88 As recently shown in healthy humans, double stimulation to different skin sites is indeed integrated, most
89 likely at spinal level, as shown by increased NWR magnitudes for double stimulation compared to single
90 stimulation (24). Moreover, the inter-electrode-distance modulated the magnitude of the NWR, suggesting
91 that spinal cord neurons do integrate spatial information (24). The use of double stimulation in the same or
92 different sites of the sole of the foot, and with varying inter-stimulus-intervals (ISIs), may provide valuable
93 information about spinal integration of nociception. The temporal discrimination (TD) of two sequential
94 stimuli, can also be used to investigate how nociceptive temporal information is processed in the
95 somatosensory system (25–29). TD likely depends on multiple stages of integration at spinal and supraspinal
96 levels (28, 30). TD for innocuous and noxious stimuli have been reported using stimuli of different nature
97 (laser, electricity and mechanical). Although TD may be representing purely temporal processes of
98 integration, it is unknown if TD also incorporates spatial information in the perception (26).

99 The present study aimed at assessing the modulatory effect of a temporal delay between two stimuli
100 delivered to the same or to two different sites in the sole of the foot in healthy humans. By using
101 simultaneous and sequential electrical stimulation with different Inter-Stimulus Intervals (ISIs), tempo-spatial
102 integration of nociceptive information in the spinal pathways was studied. It was hypothesized that shorter
103 ISIs are processed as sensory input with higher potential to produce damage and therefore associated with
104 a facilitation of the NWR. Spatial summation of the NWR when using double simultaneous stimulation was
105 also expected. The integration was primarily probed by estimating the magnitude of the NWR and
106 psychophysical measures were obtained as secondary outcomes. Temporal discrimination thresholds (TDT)
107 of the nociceptive system were also examined as an exploratory outcome.

108

109 **METHODS**

110 **Participants**

111 Fifteen healthy human subjects (9 men and 6 women, between 22 and 31 years old) participated in the study.
112 Prior to the experimental session, oral and written information was given to participants. If the subjects
113 agreed to participate, written informed consents were obtained prior to the experiment. Exclusion criteria
114 included any use of medication with known effects on the CNS that could influence the results of the study.
115 In addition, volunteers were prevented to participate in case of pregnancy or breastfeeding, previous mental,
116 musculoskeletal or nervous disorders, presence of chronic or acute pain, skin wounds on the site of
117 stimulation, or inability to cooperate during the experimental session.

118 The study was approved by the local Ethics Committee (North Denmark Region; VN-20180047) and
119 conducted in compliance with the Declaration of Helsinki.

120 **Electrical Stimulation**

121 To elicit the NWR, electrical stimulation was applied in the sole of the right foot. The skin was prepared by
122 manual abrasion of the sole of the foot to reduce skin impedance due to the thick stratum corneum. Two
123 stimulation electrodes (Neuroline 700, Ambu A/S, Denmark, reduced to a diameter of 6 mm (24, 31)) were
124 mounted on the skin over the first and fifth metatarsal bone in the sole of the foot (medially (M) and laterally
125 (L) located, respectively; Figure 1A). A large anode (7.5x10 cm; Pals, Axelgaard Ltd., Fallbrook, California, USA)
126 was placed on the dorsum of the foot for the stimulation to be perceived in the sole of the foot (31).

127 -Please, insert Figure 1-

128 Three types of stimuli were delivered: single, simultaneous, and sequential. Single stimulation through either
129 electrode (M or L), simultaneous stimulation through both M and L electrodes, and sequential stimulation
130 (through the same electrode: M or L; and as a combination of both: M, then L). Sequential stimulation was
131 delivered with varying ISIs: 30 ms, 50 ms, 100 ms, 150 ms, 200 ms and 500 ms (2). Thus, in total 21 different
132 configurations of stimuli were delivered during the experimental session. An overall interstimulus rate of 20-
133 30 s (randomized) was used to reduce habituation throughout the experiment (32).

134 -Please, insert Figure 2-

135 Two constant current computer-controlled electrical stimulators (Noxitest IES 230, Aalborg University,
136 Denmark), were used to stimulate each electrode (M and L), allowing individual control of the stimulation
137 intensity (St_i) in each electrode. St_i was individually set for M and L based on the pain threshold (P_{th}) and
138 the NWR threshold (NWR_{th}), and St_i was defined as the lowest intensity that simultaneously satisfied the
139 following conditions:

$$140 \quad St_i > 1.2 \times P_{th} \text{ AND } St_i > 1.2 \times NWR_{th}. \quad \text{Equation 1}$$

141 A staircase procedure was performed to estimate P_{th} and NWR_{th} for each stimulation electrode, following
142 an automated procedure described in a previous study (33). The criterion for P_{th} estimation was a rating of

143 perceived pain intensity above threshold ($NRS=5$) and for the NWR_{th} , a detection of a NWR when the interval
144 peak z-score of the recorded signal over the reflex windows exceeded a value of 12 (34).

145 The initial stimulation intensity was 1 mA and it was increased in steps of 2 mA until the criteria for the
146 estimation was detected (Equation 1). Subsequently, intensity was decreased by steps of 1 mA / 0.5 mA
147 (NWR_{th}/P_{th}) until the criteria was no longer met. Increasing and decreasing steps of 0.5 mA / 0.1 mA
148 (NWR_{th}/P_{th}) were then used and the thresholds were calculated averaging the last three peaks and troughs
149 derived from this process. The experiment was terminated if stimulation current reached 50 mA before the
150 criteria above (Equation 1) was met.

151

152 **Electromyography**

153 Surface electromyography of Tibialis Anterior (TA) and Biceps Femoris (BF) muscles was recorded on the right
154 leg (Figure 1B). The skin was shaved to remove hair growth and reduce impedance due to the stratum
155 corneum. Double differential configuration (35) was used by mounting three recording electrodes (Neuroline
156 720, Ambu A/S, Denmark) following SENIAM recommendations on electrode location for TA and BF (36). A
157 common reference electrode (Neuroline 720, Ambu A/S, Denmark) was placed at the patella of the ipsilateral
158 knee. Recordings were amplified, bandpass-filtered (5-500 Hz), sampled (2 kHz), and stored for offline
159 analysis.

160 **Experimental protocol**

161 Subjects were lying supine on a reclined bed with their legs relaxed. The experimental session started with a
162 familiarization protocol in which a series of stimuli were delivered in both stimulation sites (M and L) in
163 random order with varying intensities. The aim of the familiarization block was to introduce the participant
164 to the electrical stimuli and the rating tasks, and to reduce potential effects of anxiety and arousal. In the
165 second block P_{th} and NWR_{th} were estimated as previously described. During the third block, single
166 stimulation, simultaneous stimulation, and sequential stimulations were delivered (Figure 2) with five

167 repetitions of each condition in random order (21 conditions, 105 stimulations in total). Rest periods of 3-5
168 minutes duration were taken after every 25 stimulations. After each stimulation in block three, subjects were
169 asked to report an overall rating of the intensity of the perception, using a Numerical Rating Scale (NRS)
170 anchored at 0 with the label “No perception”, at 5 with “Pain threshold”, and at 10 with “Worst pain
171 imaginable”, and to indicate the stimulated site(s).

172 **Data analyses and statistics**

173 Magnitude of the NWR

174 EMG signals were filtered offline (5-500 Hz) and visually inspected to detect possible artifacts and outliers.
175 The root mean square (RMS) was calculated in the reflex window defined as 80-150ms post stimulus (2)
176 (Figure 2). Signals were then averaged across the five repetitions for each condition. Recordings were
177 excluded in case of abnormally high amplitude of both TA and BF signals, based on a robust estimation
178 method using a MAD-Median rule (37).

179 The averaged NWR due to sequential stimulation in the same electrode (M or L) was then normalized by
180 dividing it with the averaged NWR due to single stimulation in the respective electrode (M or L). For
181 simultaneous stimulation, the average of the responses to single stimulation in both sites (M and L) was used
182 to normalize. Therefore, the reported values (nNWR) express as a ratio between the NWR under a specific
183 stimulation condition (simultaneous or sequential) and the corresponding response due to single stimulation
184 in the respective site(s).

185 Temporal discrimination threshold (TDT)

186 To calculate the TDT, responses were classified as being perceived as a single or double stimulus. Stimuli
187 perceived as single were coded as “0” while stimuli perceived as double were coded as “1”. Then, the
188 responses due to the five repetitions of the same stimulus were averaged. A logistic regression was used to
189 estimate TDT (see Equation 2 below) (Frahm et al., 2018; Mørch et al., 2010; Schlereth et al., 2001).

190 Parameter “b” indicates the ISI for which participants were able to discriminate the two stimuli ($y=0.5$ in
191 Equation 2), defined as the Temporal Discrimination Threshold (TDT). The parameter ‘a’ indicates the slope
192 of the sigmoidal curve at $x = 'b'$.

$$193 \quad y = \frac{1}{1 + e^{(a(b-x))}} \quad \text{Equation 2}$$

194

195 Statistics

196 Normality of data was previously assessed in all data sets (Shapiro-Wilk test) to choose between parametric
197 or non-parametric statistical tests. Repeated measures analyses of variance (RM-ANOVA) was used when
198 normality and sphericity (Mauchly test) was confirmed, otherwise, Friedman’s test was used.

199 Friedman’s test was used to assess the effect of the ISI on the magnitude of the NWR. As differences between
200 adjacent ISIs were not of interest according to the hypotheses, post hoc analyses focused on the comparison
201 of the two extreme conditions on both sides (30 ms, 50 ms vs 200 ms, 500 ms). Planned paired comparisons
202 were conducted accordingly.

203 To assess the presence of spatial summation, RM-ANOVA was performed for pain intensity ratings during
204 single stimulation in the medial sole of the foot (M), in the lateral sole of the foot (L), and double simultaneous
205 stimulation in both sites (M and L). NRS ratings were averaged across all five repetitions of each condition
206 before conducting the statistical tests.

207 To compare the TDTs based on the spatial characteristics of the stimuli (delivered in M, L or both sites), the
208 95 % confidence intervals of the estimated thresholds were analyzed to assess significant differences.

209 When significant main effects were found, posthoc analysis with adjusted multiple comparison was
210 conducted. For non-parametric data, Wilcoxon signed rank test was performed with Bonferroni-Holm

adjustment for multiple comparison. When using RM-ANOVA, Bonferroni-Holm posthoc analysis was preferred to adjust for multiple comparisons. P-values smaller than 0.05 were considered significant.

RESULTS

Magnitude of the NWR

Compared to single stimulation, simultaneous stimulus provoked significantly larger reflexes in both TA and BF, suggesting the presence of spatial summation in the NWR (Figure 3, \emptyset : $p < 0.01$). Simultaneous stimulation elicited larger reflexes than sequential stimulation in TA (Figure 3, *: $p < 0.05$), no statistically significant difference was found in BF (see raw EMG traces of a representative subject in Figure 1C). On the other hand, sequential stimulation (averaged across all ISIs) only elicited larger reflexes in the BF muscle.

-Please, insert Figure 3-

NWR for sequential stimulation with varying ISIs

During sequential stimulation, a main effect of ISI on the TA-NWR was found regardless of the stimulated site (M: Figure 4 top row, left column $p < 0.001$; L: Figure 4 top row, center column $p < 0.01$ and the combination M-L: Figure 4 top row, right column $p < 0.05$). Stimulation with longer ISIs elicited larger reflexes in TA, regardless of the stimulated site (M, L, and their combination; Figure 4 top row * $p < 0.01$).

For the BF muscle, a significant main effect of ISI was found when stimulating through the L electrode (Figure 4 bottom row, center column $p < 0.05$) and in the combination of both M and L (Figure 4 bottom row, right column $p < 0.001$). When stimulating through the M electrode, BF-NWR showed a similar tendency towards smaller reflexes for longer ISIs (Figure 4 bottom row, left column $p = 0.06$). Results of post hoc analysis showed that longer ISIs produced decreasing NWR magnitudes in BF (Figure 4 bottom row * $p < 0.05$).

233

234 -Please, insert Figure 4-

235

236 **Perceived intensity**

237 The mean perceived intensity responses to all stimuli were perceived above pain threshold (NRS=5),
238 regardless of stimulated site or condition (single, simultaneous, or sequential).

239 As shown in Figure 5, the pain intensity following sequential and simultaneous stimulation were perceived
240 as more painful than single stimulation ($p < 0.05$, Wilcoxon signed rank test), however, no difference was
241 found between simultaneous and sequential stimulation.

242 -Please, insert Figure 5-

243 For sequential stimulation with different ISIs, regardless of the electrode site, there was no significant effect
244 of the ISI ($p > 0.05$, Friedman's test) (Figure 6).

245 -Please, insert Figure 6-

246

247 **Temporal discrimination thresholds**

248 The estimated temporal discrimination thresholds (TDTs) are summarized in Table 1. The TDTs were 84.1 ms
249 for the sequential stimulation in the medial site (M), 95.5 ms for stimulation in the lateral electrode (L); and
250 71.0 ms for stimulation as a combination of both electrode sites (M-L). The comparison between conditions
251 showed that significant lower TDT was observed when stimulating both sites (M-L), rather than any single
252 site. Additionally, TDT was lower for stimulation in the medial electrode (M) compared to the lateral
253 electrode (L).

254

255

256 *Table 1: Results from the logistic curve fitting to Equation 2. Results expressed as mean of each stimulation type with 95%*
257 *Confidence intervals of the logistic regression.*

Stimulation Type	TDT [ms]	95% CI [ms]
Medial electrode (M)	84.1	80.9-87.1
Lateral electrode (L)	95.5	93.5-97.6
Combination (M-L)	71.0	69.4-74.2

258

259 **DISCUSSION**

260 The current study investigated the tempo-spatial integration of double stimulation in healthy humans by
261 assessing the magnitude of the NWR in one proximal and one distal muscle (BF and TA, respectively) during
262 different stimulation conditions. Spatial summation was found in both spinal- (BF-NWR, TA-NWR) and
263 supraspinal- (perceived intensities) mediated outcomes. Sequential stimulation with varying ISIs (between
264 30-500 ms) had a significant modulatory effect on the magnitude of the NWR but not on the perceived
265 intensities. Interestingly, the modulation affected proximal (TA) and distal (BF) muscles differentially. For
266 increasing ISIs, the TA-NWR were facilitated, while the opposite pattern was seen in the BF muscle. The
267 temporal discrimination threshold was lower for stimuli delivered at different sites, compared to a single
268 stimulation site.

269 **Tempo-spatial integration in spinal nociception assessed via the NWR**

270 Simultaneous double stimulation elicited a significantly larger NWR than single stimulation in both TA and BF
271 suggesting that the spatial characteristics of the stimulation modulates the magnitude of the NWR (EMG
272 traces of a representative subject in Figure 1C). When using simultaneous stimulation, the elicited NWR had
273 an average magnitude of 150 % of the single stimuli (Figure 3). This observation is likely reflecting sub-additive
274 spatial summation (SS) of the NWR in healthy humans. Evidence regarding SS in the somatosensory system
275 is abundant in the literature, particularly in studies that used the intensity of the perception as an outcome
276 of sensory integration (38–47). The NWR, being a polysynaptic reflex that reflects spinal nociceptive
277 processing (48, 49), has not been explored for concurrent stimuli at different locations, except for one
278 previous study (24). Our previous study and the current study show that simultaneous stimulation at two
279 sites located medially and laterally in the sole of the foot (Figure 1A) elicits a NWR approximately 1.5 times
280 larger than when stimulating the individual sites. These results support the notion that spinal cord neurons
281 integrate afferent information from nociceptive fibers innervating skin regions in different dermatomes (40,

282 50, 51). From a defensive point of view, it is crucial to exploit this information since it allows an adaption of
283 the reflex magnitude according to the spatial characteristics of the stimuli.

284 The difference in the NWR amplitude across varying ISIs, show that the temporal characteristics of the stimuli
285 also play an important role in the modulation of the NWR (Figure 4). With increasing ISIs, the magnitude of
286 the TA-NWR increased, while the BF-NWR decreased. Interestingly, this modulation seems to be independent
287 of the stimulation site since the same tendencies were observed during stimulation of the medial electrode
288 (M), the lateral electrode (L), and with the combination of electrodes (M-L) (Figure 4).

289 One possible explanation for the TA facilitation for larger ISIs (Figure 4; top row), could be varying degrees of
290 temporal summation (TS) modulating the magnitude of the reflex. Modulation of the NWR due to TS, has
291 previously been demonstrated (16, 17, 20, 52), however, the current study is the first to investigate TS in
292 proximal and distal muscles concurrently when delivering stimulations in two different locations with
293 different ISIs. A few human studies have compared the effect of the stimulus ISI on TS (16, 19, 52). Those
294 studies showed that shorter ISIs generally produce larger TS effect. Although those observations seem to
295 contradict the present ones, it is worth noting that the assessment of the magnitude of the NWR in previous
296 studies was solely performed on proximal muscles: Semitendinosus (52), BF (16, 19) and Rectus Femoris
297 muscles (19). Thus, those prior findings are consistent with the present results regarding the BF-NWR
298 modulation. Comparing the modulation of TA-NWR and BF-NWR, it appears that the NWR modulation for
299 varying ISIs differs in proximal and distal muscles consistently across stimulation sites. Thus, it seems
300 plausible that the differential modulation on TA and BF reflect the protective role of the NWR. A sequential
301 stimulation with shorter ISI is likely associated with afferent input with higher potential for damaging the
302 body, regardless of the specific stimulated site on the foot. The optimal motor response to a more dangerous
303 stimulus would be to generate a strong withdrawal of the entire limb, by preferentially recruiting proximal
304 muscles. Stimuli with longer ISIs, on the other hand, do not seem to trigger such facilitated reflex magnitudes

305 in the proximal muscles, suggesting that the recruitment of distal muscles (TA) could produce a sufficient
306 withdrawal through dorsiflexion of the foot.

307 Interestingly, the temporal discrimination thresholds, suggest that stimuli with ISIs shorter than 70 ms are
308 discriminated correctly in less than 50 % of all cases, regardless of the site being stimulated (see TDT in Table
309 1). In these cases, a crude withdrawal of the limb (by contraction of proximal muscles, e.g.: BF) seems to be
310 the safest and optimal behavioral response to that noxious threat, rather than a more refined foot
311 withdrawal (TA) (Figure 4).

312 In TA it was found that sequential stimuli produced significantly smaller reflexes compared to simultaneous
313 stimuli (Figure 3). The spatial summation phenomenon that integrates simultaneous stimulation of M and L
314 seems to diminish when the simultaneity is lost. This observation further argues for a dynamic tempo-spatial
315 integration of nociception present in the spinal cord circuitry governing the NWR (2, 6, 11).

316 In summary, it appears that when delivering sequential stimulation, the nociceptive afferent volley is
317 conveying temporal information, which is integrated within spinal cord reflex circuitry to elicit an optimal
318 reflex response. With an ISIs short enough to prevent sensory discrimination as independent stimuli, the
319 NWR circuitry seems to be responding as if the limb was exposed to a bigger threat (Figure 4) facilitating
320 recruitment of proximal muscles (BF).

321 **Descending modulation of the NWR**

322 Evidence of supraspinal structures modulating motor defensive behavior have previously been reported (53–
323 56). Specifically, the midbrain periaqueductal gray (PAG) likely play an important role in sensorimotor
324 integration and exerts modulation over the motor system to assist survival (57). In the present study,
325 particularly when using long ISIs (200-500 ms), the time between the first and second stimulus is sufficiently
326 large to see top-down modulation of the NWR (58) triggered by the first stimulus (the NWR is reported as
327 the size of the last NWR when using sequential stimulation, see Figure 2).

328 Moreover, it has been shown that different emotional states can modulate the magnitude of the NWR (59–
329 62), which is outlining a pathway from cognitive processes to up/down regulation of the excitability in the
330 spinal reflex pathways. Although preventive measures were taken in this study to familiarize the participants
331 to the stimulation protocol, a top-down modulation of the NWR ascribed to the cognitive state cannot be
332 completely discarded. Experiments involving painful stimulation, may induce a state of anxiety that can
333 modulate spinal nociception (62). Evidence has recently been reported showing facilitation of the TA-NWR
334 when the subject is exposed to an anxiety induced stimulation paradigm involving heterotopic painful
335 electrical stimulation (62). Thus, it is interesting that the magnitude of TA-NWR was significantly larger for
336 larger ISIs, which might involve a transcortical loop. However, if anxiety affected the subjects, the facilitation
337 of spinal nociception would most likely affect the magnitude of the NWR for all ISIs. Additionally, the
338 magnitude of the BF-NWR was significantly reduced with longer ISIs, suggesting that the modulatory effect
339 of ISIs on the NWR magnitudes cannot be explained by changes in the cognitive state exerting spinal
340 modulation.

341 The subjects' attention to the stimulus may induce inhibition of the NWR (63). Hence, in the present study,
342 the first stimulus may draw the attention of the subject leading to smaller reflexes with the second stimulus.
343 However, since stimuli were totally randomized and subjects were instructed to identify stimulated site in
344 every trial, it seems unlikely that this effect is restricted only to some ISIs.

345 It is not known to what extent descending modulation differentially modulate proximal vs distal muscles. A
346 recent study, however, has suggested that the unpredictability of the onset of an imminent stimulus
347 produced differential modulation of proximal vs distal muscles involved in the NWR (64). In that study, when
348 the onset of a second incoming stimulus (with an ISI of 1 s) was unknown (as in our study), the size of the
349 NWR was differentially modulated with larger TA-NWR in combination with smaller BF-NWR. This is in
350 agreement with the present results since the first stimulus may serve as a cue, particularly in the conditions

351 with larger ISIs. This further support the NWR as a dynamic and highly adaptable behavioral response that
352 integrates temporal and spatial information of the stimulus to produce optimal withdrawal.

353 **Perceived intensities**

354 Higher perceived intensity ratings were found for simultaneous and sequential compared to single
355 stimulation, most likely caused by spatial summation (Figure 5). Although the exact mechanisms behind the
356 summation phenomenon remains to be clarified, previous findings in animal studies and indirect evidence in
357 human studies have shown that deep dorsal horn neurons likely play a major role in this central integration
358 (38–40, 42, 65). From the present results, the fact that stimulated areas on the medial and lateral parts of
359 the sole of the foot belong to different dermatomes (66, 67), likely support central neural circuits being
360 responsible of the observed spatial summation in the perceived intensities.

361 However, it is important to notice, that although statistically significant, the average difference between
362 simultaneous (and sequential) versus single stimulation in the perceived intensity of pain was less than half
363 a point on the NRS, a similar difference in pain intensity as reported in other studies assessing SSP (40, 43,
364 46).

365 Contrary to the NWR modulation previously discussed, different ISIs did not modulate the perceived
366 intensities (Figure 6). If shorter ISIs were indeed encoded as sensory input with more potential to produce
367 damage, larger perceived intensity ratings may also be expected. However, this was not observed in this
368 study (Figure 6). One reason that could explain this apparent disagreement between the reflex behavior and
369 the intensity of the perception is that a larger number of stimuli are needed to establish a significant
370 modulation in the perceived intensity due to temporal summation (68). Indeed, a previous study on temporal
371 summation of pain and spinal nociception have shown that a significant modulation of pain perception is
372 achieved when the skin is repetitively stimulated only after the delivery of the third stimulus with an ISI of
373 500 ms (69). The NWR responses observed in this study is affected by temporal summation which is already

374 seen following the second stimulus (17), suggesting a larger reflex sensitivity to repetitive stimulation, while
375 a larger number of pulses is needed to produce a modulation of the perceived intensity.

376 **Temporal discrimination**

377 Previous observations differ regarding the tempo-spatial discriminatory capabilities of the somatosensory
378 system for noxious and innocuous stimuli. Specifically, better directional discrimination for noxious vs
379 innocuous laser stimulation has been reported (70). However, when comparing noxious laser stimuli against
380 innocuous mechanical stimulation the opposite was found (e.g. better discrimination for innocuous
381 stimulation) (71, 72). The TDT in the present study (noxious) ranged between 71.0 ms and 95.5 ms (Table 1).
382 For innocuous electrical stimulation in the lower limb, TD has been reported between 39.9 ms and 41.2 ms
383 (25). A possible reason for the disagreement between the studies is likely the stimulation modality (laser,
384 mechanical and electrical), which will affect the discrimination (73). Laser stimulation leads to a natural
385 activation of nociceptors while electrical stimuli recruit a mix of A δ and A β fibers, by bypassing the
386 transduction processes and are associated with a high level of synchronization in the afferent volleys. Thus,
387 temporal discrimination will also depend on the stimulation modality, and this may explain the disagreement
388 between previous studies and the current study.

389 The TDT depends on the stimulation sites (M and L), for these two sites it is possible that different stratum
390 corneum (SC) thicknesses explain the better improved temporal discrimination when the stimuli are applied
391 in the medial side (thinner SC). Interestingly, the inclusion of a spatial factor (i.e. combination of M and L), in
392 which the first stimulus is delivered in the medial side and the second stimulus in the lateral side of the sole
393 of the foot, significantly improved the temporal discrimination of the stimulus (Table 1), arguing that spatial
394 information is integrated and the perception reflects the temporal characteristics of nociceptive information.

395 **Limitations**

396 The NWR is a response elicited to defend the body from a potentially damaging stimulus. Together with this
397 reflex response, the perception of pain may be elicited by the stimulus. The processing of the noxious

stimulus that produces a defensive reflex response and a perceptual experience of pain involves several structures from the afferent fibers through the spinal cord and to the brain. In the present study, electrical stimulation was used to elicit the NWR, potentially depolarizing unmyelinated as well as myelinated fibers. Overall, a contribution of large myelinated fibers in the elicitation of the NWR cannot be completely discarded (74, 75). Given the duration of the recording window for the NWR, a contribution from supraspinal centers is likely. Previous studies have suggested that deep dorsal horn neurons encode spatial integration of nociception and do not project to ascending pathways (10, 50, 76). Likely arguing for spinal circuitry responsible, at least to some extent, for spatial integration and elicitation of the reflex response. On the other hand, perception involves many supraspinal structures to produce the perception of pain. A clear separation and independent assessment of both processes is, therefore, not possible in human studies.

Conclusion

The NWR circuitry is integrating the spatial characteristics of a sensory input, e.g. observed as larger reflex responses in both TA and BF when simultaneously stimulating two sites (compared to single stimulation; Figure 3). A similar phenomenon of summation was observed for perceived intensities (Figure 5). Moreover, TS of the NWR for sequential stimulation was found in both TA and BF muscles, and when the same and different sites were stimulated (Figure 4). TS was largest for small ISIs in the BF muscle, while the opposite pattern was observed in TA (larger TS for longer ISI). The perceived intensities were not modulated by varying ISIs (Figure 6). The sensory information concerning the temporal and spatial aspects of the stimulation shapes both the reflex response and the perception in terms of temporal discrimination. The different tendencies in the modulation observed in the TA and BF muscles for different ISIs and for different stimulation sites (Figure 4) suggest that spinal cord neurons are playing a role in the tempo-spatial integration of nociceptive information, leading to a reflex response which is distributed across multiple spinal cord segments and governed by an efficient defensive withdrawal.

Conflict(s) of Interest: None

422 **Source(s) of Financial Support for the Project:** This study was funded by the Danish National Research
423 Foundation (DNRF121) and by the European Union's Horizon 2020 research and innovation programme
424 under the Marie Skłodowska-Curie grant agreement No 754465.

425 **Ethical Permissions:** Written informed consents were obtained from each volunteer previous to their
426 participation. The local ethical committee approved this study (N-20180047).

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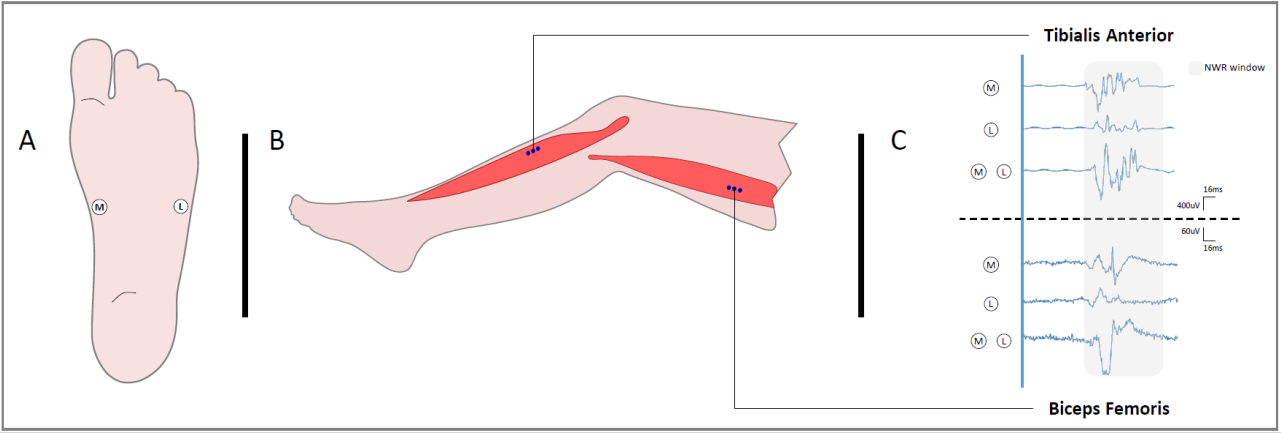
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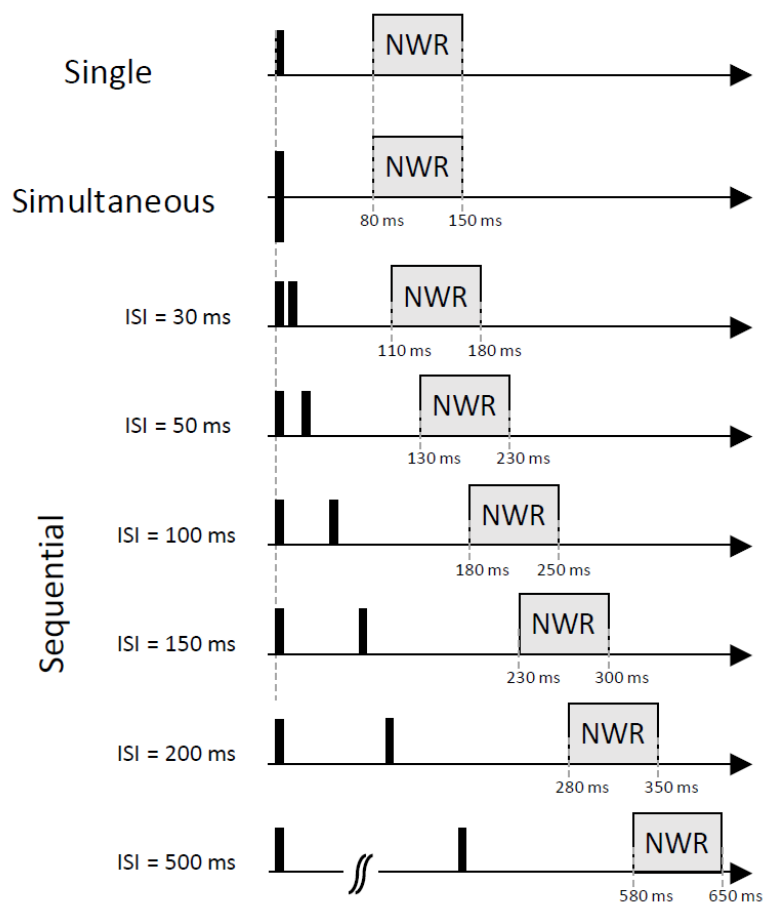
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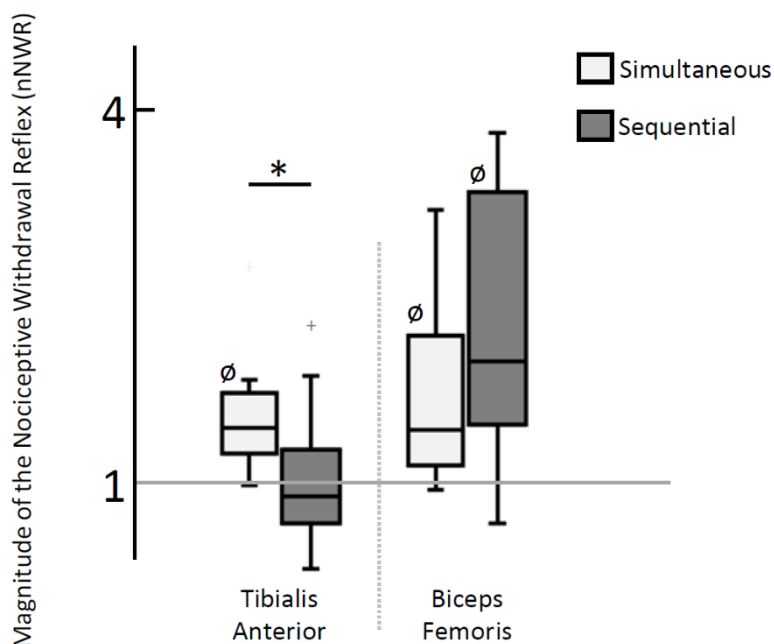
631 **Figure 1:** Diagram indicating the stimulated sites, recorded muscles and representative raw EMG traces
632 obtained with Single and Simultaneous stimulation. A) Figure showing the location of the stimulating
633 electrodes in the sole of the foot: M and L, medially and laterally located, respectively. B) Distal (Tibialis
634 Anterior) and proximal (Biceps Femoris) muscles from which C) sEMG recordings were obtained. Raw EMG
635 traces for both TA and BF illustrate the response of a representative subject to single and simultaneous
636 stimuli in M and L sites.



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638 **Figure 2:** Stimulus types used in the experiment: Single, Simultaneous and Sequential. Black rectangles
 639 indicate stimulus artifact. The NWR was quantified in a 70 ms window (blue box) starting 80 ms after the
 640 trigger of second stimulation. Sequential stimulation was delivered with varying inter-stimulus intervals
 641 ranging from 30 ms to 500 ms.

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644 **Figure 3:** Box and whiskers plot showing nNWR for simultaneous and sequential (averaged across ISIs)

645 stimulation in both TA (left) and BF (right) muscles. For TA muscle, there was a significant difference

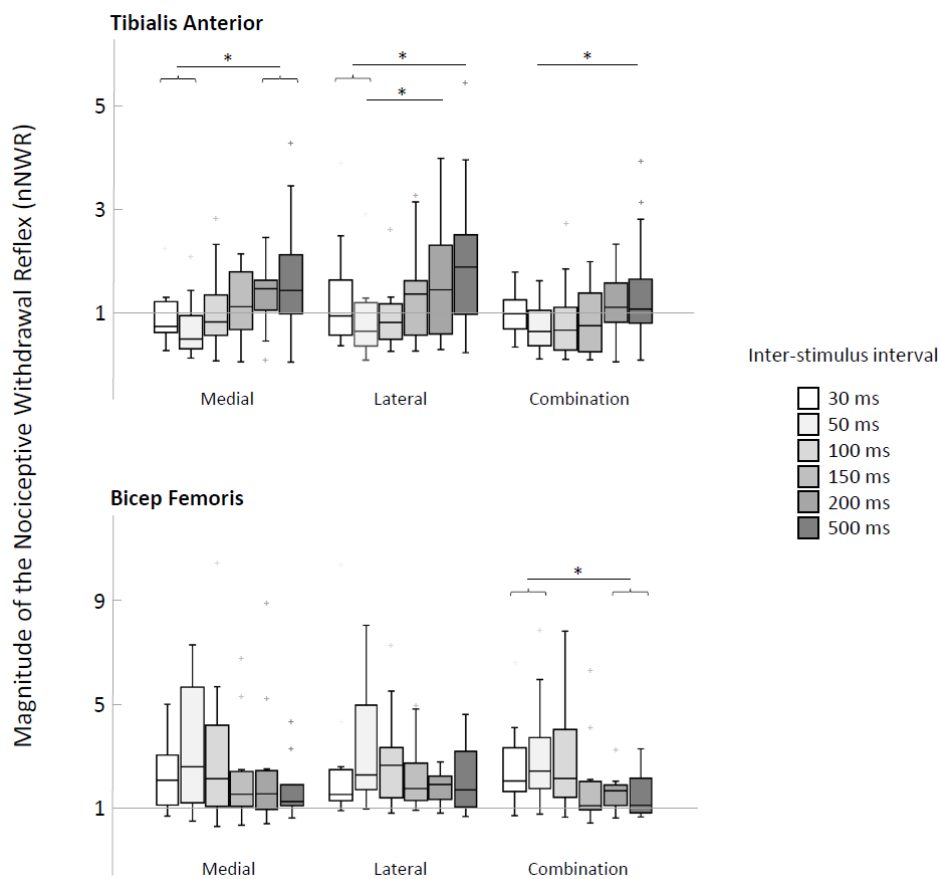
646 between simultaneous and sequential stimulation (*: $p < 0.05$, Wilcoxon signed rank test). When compared

647 to single stimulation (not shown), simultaneous stimulation showed increased reflex magnitude for both TA

648 and BF muscles (\emptyset : $p < 0.01$, Friedman's test). Sequential stimulation, however, only provoked larger

649 reflexes than single stimulation in BF (\emptyset : $p < 0.01$, Friedman's test).

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Figure 4: Box and whiskers plot illustrating the magnitude of the NWR when using sequential stimulation with different ISIs (30ms – 500ms). The top row shows the results obtained in the Tibialis Anterior muscle, while the bottom row displays the results of the Biceps Femoris muscle. ISIs. Values are shown as nNWR, normalized to the NWR due to single stimulation (see Methods section). Opposite tendencies in the proximal (BF, bottom row) compared to the distal (TA, top row) muscles were observed. For larger ISIs, TA-NWR was facilitated while BF-NWR was reduced (*: $p < 0.05$, Wilcoxon signed rank test).

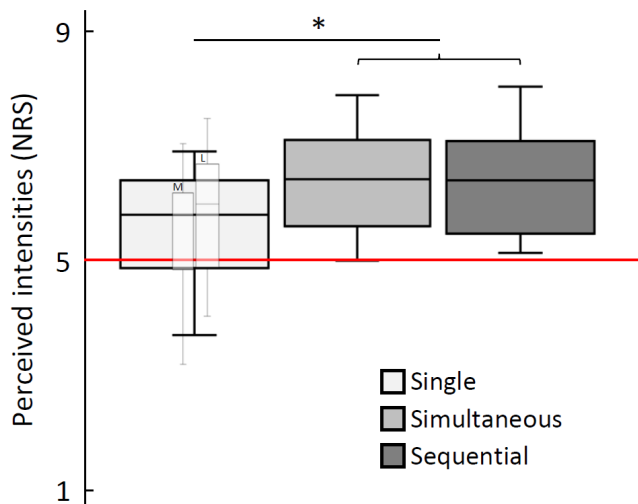


Figure 5: Box and whiskers plot showing Pain intensity ratings due to Single, Simultaneous and Sequential stimulation of M and L. Simultaneous and Sequential stimulation were perceived as more painful than Single stimulus (* $p < 0.05$, Wilcoxon signed rank test). No significant difference was found between Simultaneous and Sequential stimulus. Pain threshold (NRS=5): red horizontal line.

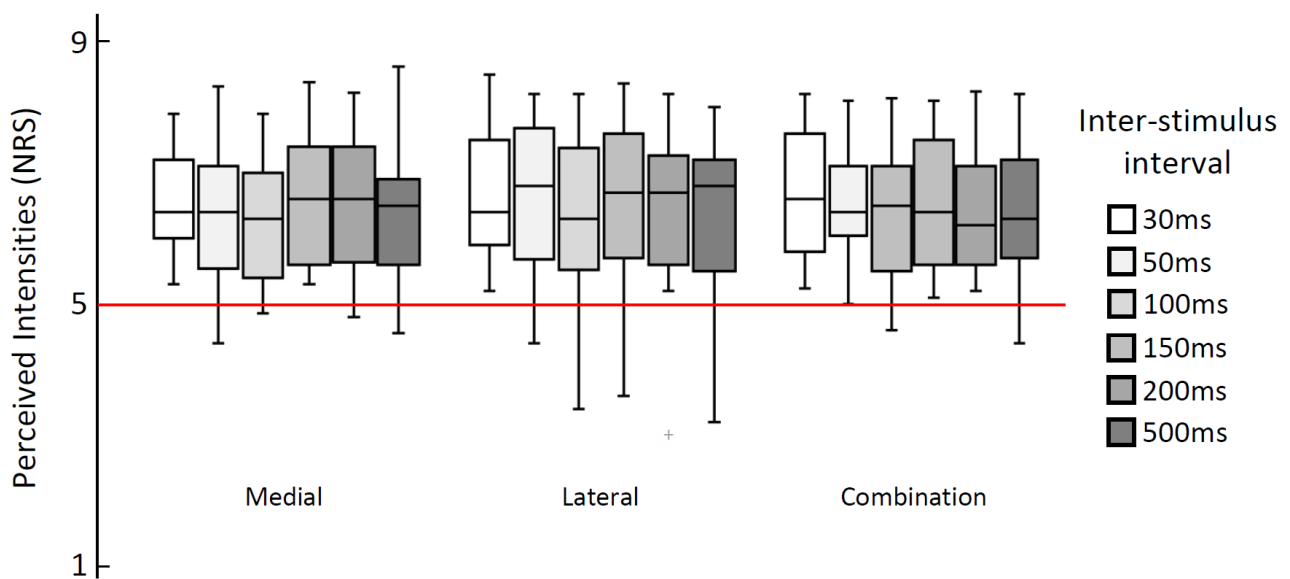


Figure 6: Box and whiskers plot of pain intensity ratings reported for Sequential stimulation in the medial electrode (M), in the lateral electrode (L) and as a combination of both (M-L). Different Inter-Stimulus Intervals (ISIs) are shown. Perception was above pain threshold (NRS=5: red horizontal line) in all cases, and there was no significant effect of ISI on the perception of pain intensity ($p > 0.05$, Friedman's test).