

Personal influencing factors for pressure pain threshold in healthy people: A systematic review and meta-analysis

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Personal influencing factors for pressure pain threshold in healthy people: A systematic review and meta-analysis



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ABSTRACT

All studies that investigated personal factors influencing pressure pain threshold (PPT) in healthy people were synthesized. Data was summarized, and risk of bias (RoB) and level of evidence were determined. Results were pooled per influencing factor, grouped by body region and included in meta-analyses. Fifty-four studies were eligible. Five had low, nine moderate, and 40 high RoB. Following meta-analyses, a strong conclusion was found for the influence of scapular position, a moderate for the influence of gender, and a weak for the influence of age (shoulder/arm region) and blood pressure on PPT. In addition, body mass index, gender (leg region), alcohol consumption and pain vigilance may not influence PPT. Based on qualitative summary, depression and menopause may not influence PPT. For other variables there was only preliminary or conflicting evidence. However, caution is advised, since the majority of included studies showed a high RoB and several were not eligible to include in meta-analyses. Heterogeneity was high in the performed meta-analyses, and most conclusions were weak. More standardized research is necessary.

1. Introduction

Pain is defined as "an unpleasant sensory and emotional experience associated with, or resembling that associated with, actual or potential tissue damage" (Raja et al., 2020), and is one of the most common reasons why people opt for medical help (Mäntyselkä et al., 2001). Pain has a huge impact on a person's functioning and quality of life, but also on society, as pain usually results in high medical costs or other socio-economic problems (Blyth et al., 2019).

The somatosensory system processes nociceptive signals that can lead to the sensation of pain (Yam et al., 2018). Psychophysical testing, such as quantitative sensory testing (QST), can be used to assess sensitivity of the somatosensory system and associated pathways, in which measuring pain thresholds is an indispensable part (Backonja et al., 2009). Based on patients' self-reported sensory experience, pain thresholds for mechanical, thermal, vibration and electrical stimuli can identify allodynia and hyperalgesia (Cruz-Almeida and Fillingim, 2014). These are regarded as signs of altered somatosensory processing (Jaber et al., 2018). In clinical practice and research, the most feasible way to assess pain thresholds is measuring pressure pain thresholds (PPTs), which refer to the minimum amount of pressure necessary to induce pain (Hall et al., 2015). A pressure sensation stimulates C- or Aδ- fibres in the skin, of which a signal goes through the anterior spinothalamic tract until it reaches the thalamus in the brain, which processes the information before sending it out to various parts of the cortex (Yam et al., 2018). To determine the PPT, usually an algometer is used, which is found to be reliable and valid (Frank et al., 2013). To date, PPT values could be a useful measure for possible signs of altered somatosensory processing, and thus to detect differences between healthy and patient populations (diagnostic) or changes over time (e.g., before and after treatment, responsive) (Jaber et al., 2018; Arendt-Nielsen and Yarnitsky, 2009; Walton et al., 2011).

In order to adequately interpret PPT values, normative values are necessary (Arendt-Nielsen et al., 2018; Vardeh et al., 2016). The first

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step is to focus on the diagnostic goal, as these normative values are the only way to assess whether a patient presents altered somatosensory processing or not. However, to date, no clear normative values for PPT are available and determining them seems to remain challenging. Some literature of normative data in healthy people exists, but even when groups based on age and measurement location were considered, a broad range of PPT values is still presented (Waller et al., 2016; Pfau et al., 2014; Sterling et al., 2002; Tuveson et al., 2006). This variation could be explained by other factors influencing PPT as well.

Systematic reviews and meta-analyses found that age (4 studies) (Tumi et al., 2017), and gender (33 (Racine et al., 2012) and 5 studies (Riley et al., 1998)) influenced PPT in healthy people. Unfortunately, other reviews, also focusing on influencing factors (such as the use of alcohol, gender role, or menstrual cycle), only reported analyses combining all pain thresholds for different modalities (Martin, 2009; Iacovides et al., 2015; Horn-Hofmann et al., 2015; Alabas et al., 2012). These modalities stimulate different fibres through different pathways: e.g. mechanical (pressure) and thermal stimuli stimulate C- or Aδ- fibres through the anterior and lateral spinothalamic tract, respectively (Yam et al., 2018). Tumi et al (Tumi et al., 2017). confirmed this by finding a difference in the influence of age between heat pain threshold and PPTs, and Riley et al. (1998) found larger gender differences for PPT compared to thermal stimuli in healthy people. Thereupon, differences in the targeted anatomical structure and tissue depth exist (Fillingim, 2002). Aforementioned differences could lead to different responses to pain threshold testing, and therefore it is important to analyse a certain pain threshold modality separately. As such, the influence of other personal variables (other than age and gender) on PPT specifically remains unclear. Moreover, the last review of the influence of gender on PPT dated from 2012 (Racine et al., 2012), so an update seems necessary as well.

To date, no guidelines are available in scientific literature for which influencing factors to consider when determining normative values for PPT. It is important to know which factors influence the PPT, because this information can be useful for the diagnosis of patients with altered central somatosensory processing. These normative values can thus be used to detect an indication of the presence of mechanical hyperalgesia. As such, more research to detect the influence of different factors on PPT separately is necessary. This review will focus on all personal clinically measurable influencing factors, because measurements can be standardized regarding environmental factors and time of measurement throughout the day, but also because invasive and medical lab tests (e.g., to test the influence of genetics, fat mass) are not always available in clinical practice (such as physiotherapists cabinet).

Therefore, the aim of this systematic review was to synthesize all studies that had the purpose to explore which clinically measurable personal factors might influence PPTs in healthy people. This way, these personal factors can be considered in future studies on normative values of PPT, and can be considered for diagnosis of patients with mechanical hyperalgesia.

2. Methods

This systematic review and meta-analysis was conducted according to the updated Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines (Page et al., 2021).

2.1. Eligibility criteria

Studies were included if they evaluated clinically measurable personal factors (I) possibly influencing PPT (O) in a healthy population (P). Personal influencing factors were defined as moderators, predictors or correlating personal characteristics for PPT values, or as mean difference PPT values between groups divided based on the personal factor. The eligibility criteria can be found in Table 1.

Table 1

Eligibility criteria related to PICO.

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	Inclusion	Exclusion
Р	 Healthy human subjects free of pain, illness or disease (included in intervention or control group) > 18 years of age 	1. Animal studies
I	 Clinical, non-invasive measurable personal factors influencing the PPT (e.g., gender, strength, psy- chosocial factors) 	 Other factors such as environmental factors (e.g., attention, type of assessor)/ genetics
С	 No comparison with another population, comparison with a non-healthy population or com- parison between two healthy populations (e.g. men vs women) Separate statistical analyses for 	 Statistical analyses only of mixed population (e.g., patient population and healthy subjects)
0	the healthy subjects 1. PPT measured with pressure algometer or other material that can measure the amount of pressure objective	 Vibration, electrical, thermal, ischemic pain threshold Vibration, electrical, thermal, sensory, ischemic, mechanical detection threshold
S	 All sorts of study designs in a longitudinal setting 	 Reviews, Meta-analyses, Abstracts, Letters, Congress proceedings, case reports, cross-sectional or case- control studies (PPT measured on only one time point)
L	1. Articles written in English, Dutch, German or French	1. Articles written in any other language

Abbreviations: P, population; I, intervention; C, comparison; O, outcome; S, study design; L, language

2.2. Information sources and search strategy

The electronic databases PubMed (MEDLINE), Web of Science (WoS) and Embase were searched for eligible literature up to 11th January, 2022. Additionally, the references of previous reviews (Tumi et al., 2017; Martin, 2009; Horn-Hofmann et al., 2015; Alabas et al., 2012) and the references of the included articles retrieved through the search strategy were screened for relevant studies. To answer the research question, three different sets of key words (P, I and O) were composed following the Patient (P), Intervention (I), Comparison (C), Outcome (O) and Study design (S) model and combined using 'AND' and 'OR' (Eriksen and Frandsen, 2018). The search strategy for PubMed can be found in Table 2, the strategies for WoS and Embase can be found in table S1. No additional search filters were added.

2.3. Selection process

All studies retrieved from the electronic databases were imported in Endnote 20 (\$author1\$ et al., 28. </id>

Table 2	
Search Strategy related to PICO in PubMed.	

Population	Intervention	Outcome
("Healthy Volunteers"[Mesh]) OR Healthy voluntee* OR healthy people OR healthy subjec* OR healthy perso* OR healthy individua*	("Prognosis"[MeSH Terms] OR "effect modifier, epidemiologic"[MeSH Terms]) OR Predict* OR moderat* OR modif* OR Prognos* OR "epidemiologic effect modifier" OR influenc*	("Pain Threshold"[Mesh] OR "Pain Measurement"[Mesh] OR "Pain Perception"[Mesh]) OR "pain threshold" OR "pain measurement" OR "pain perception" OR "quantitative sensory testing" OR qst OR pressure algomet* OR "mechanical pain threshold" OR pressure pain threshold OR ppt

Abbreviations: QST, Quantitative Sensory Testing

Table 3

Level of evidence and strength of recommendation scoring.

	Level of evidence		Strength of recommendation
LoE 1 *	Systematic review of randomized trials	SoR I (very strong)	At least one LoE 1 study or three LoE 2 studies
LoE 2 *	Randomized trial	SoR II (strong)	At least one LoE 2 study or three LoE 3 studies
LoE 3 *	Non-randomized controlled cohort/follow-up study	SoR III (moderate)	At least one LoE 3 study or three LoE 4 studies
LoE 4 *	Case-series, case-control, or historically controlled studies	SoR IV (weak)	At least one LoE 4 study or three LoE 5 studies
LoE 5 *	Mechanism-based reasoning	SoR V (N/A)	At least one systematic review of descriptive and qualitative studies
	ũ	SoR VI (N/A)	At least a single descriptive or qualitative study
		SoR VII (very weak)	At least one LoE 5 study
		Preliminary SoR	Based on only one study
		Conflicting conclusion	Conflicting results

Abbreviations: RCT, Randomized Controlled trial; LoE, Level of Evidence; SoR, Strength of recommendation; N/A, not applicable *Level of evidence can be graded down due to study quality or other methodological issues.

EndNote Team</au></Author></AuthGrp>) and identified duplicates were removed. The remaining studies were independently screened on title and abstract by two reviewers (SV and VH) with the help of Rayyan (Ouzzani et al., 2016). Subsequently, potentially eligible studies were additionally screened on full text by both reviewers independently in the following order of exclusion: language > study design > outcome > population > intervention. Conflicts during the whole process were resolved by consensus and in case of doubt, the last author was contacted.

2.4. Data collection and items

Data of all included studies were extracted into an evidence table. Information about (Raja et al., 2020) Author, year and origin; (Mäntyselkä et al., 2001) Study design; (Blyth et al., 2019) Participants, such as group composition and characteristics, and eligibility criteria; (Yam et al., 2018) Device, speed of the pressure build-up and analysis of PPT, including the reported signal and patient position; (Backonja et al., 2009) Location of PPT; (Cruz-Almeida and Fillingim, 2014) Influencing factor and measurement method; and (Jaber et al., 2018) Results was collected. The first reviewer (SV) completed the evidence table and the second reviewer (VH) checked the table independently.

2.5. Risk of bias in individual studies

Risk of bias (RoB) in the individual studies was assessed using the quality in prognostic studies (QUIPS) checklist (Hayden et al., 2013), as the aim of our review was to find the prognostic factors for PPT. The checklist consists of six domains that can be scored either as a 'high risk', 'moderate risk' or 'low risk' of bias: 1) Study Participation, 2) Study Attrition, 3) Prognostic Factor Measurement, 4) Outcome Measurement, 5) Study Confounding, and 6) Statistical Analysis and Reporting. Risk of bias assessment was performed independently by two reviewers (SV and VH), and conflicts were resolved by consensus. To assure uniform RoB scoring, guidelines for interpretation of each item were discussed beforehand through a calibration exercise. The overall RoB judgement of a study was based on all domains; ranging from overall 'low' RoB if all domains were scored 'low' or maximum one 'moderate'; to an overall 'high' RoB if at least one domain was scored as 'high' or \geq 3 as 'moderate'. All other studies were judged as having an overall 'moderate' RoB.

Additionally, the overall level of evidence per study was evaluated by the first author (SV) based on RoB score and study design with the Centre for Evidence-Based Medicine (CEBM) guidelines (Explanation of the, 2011). Different level of evidence (LoE) scores were given based on their study design, methodology and RoB score (Table 3).

The RoB scoring was completed independently and in a double-blind manner by two reviewers (SV and VH). Finally, results were compared and conflicts were resolved by consensus or by contacting the last author. Afterwards, the first author bundled all the results per personal influencing factor on PPT and strengths of recommendations were made, divided into different categories based on the CEBM guideline (Table 3)

(Ackley, 2008).

2.6. Statistical synthesis methods and effect measures

Studies that presented mean and standard deviation for PPT values were combined and presented in different forest plots per influencing factor (grouped by body region) with the software Review Manager (RevMan) 5.4.1 (Review Manager Web, 2020), as subgroup analyses is possible with this software. For studies that presented correlation coefficients the software Jamovi 1.6.23 (The jamovi project, 2021) was used, as RevMan does not provide this correlation meta-analysis function. A Fisher r- to z- transformed correlation coefficient was used in this case. Only studies with full available data were included in the analysis, and authors of the original studies were contacted if data was missing. Analysis went further when the authors did not respond for a period longer than four weeks. Data was pooled through calculating the mean value if multiple categories of the influencing factor were present (e.g., absolute values and correlation coefficients of PPT values [in case of univariate analyses] from women and men were pooled to get a clear view of the influence of the factor age and vice versa, data of different locations was pooled to one data per body region). Standardized mean difference (SMD) was used to compare data, if different units for measuring PPT were used in the included studies. The presence of heterogeneity was assessed with the value I². In case of high heterogeneity $(I^2 > 50\%)$, random effects methods were used, and in case of low heterogeneity ($I^2 < 50\%$) fixed effects methods were used. Subgroup analyses per body region were performed if possible. Small-study effects were checked by visual observation of the symmetry of a funnel plot, but could only be checked if more than 10 studies were implemented in the meta-analysis (Debray et al., 2018; Sterne et al., 2011). Afterwards, sensitivity analyses correcting for studies that did not report full eligibility criteria and correcting for the model of meta-analyses (random or fixed effects) were performed as well. An overall p-value of the models was obtained in which significant results meant p < 0.05. An effect size with an overall SMD < 0.2 was considered very small, 0.2–0.5 small, 0.5–0.8 medium, and > 0.8 large (Andrade, 2020). An overall correlation coefficient (CC) < 0.2 was considered very weak, 0.2–0.39 weak, 0.4-0.6 moderate, 0.6-0.79 strong, and 0.8-1.0 very strong (11. Correlation and regression, 2021).

3. Results

3.1. Study selection and characteristics

The study selection process is illustrated in the PRISMA flowchart (Page et al., 2021) (Fig. 1). Our search strategy resulted in 32 eligible studies (Alfieri et al., 2017; Alves et al., 2017; Andrzejewski et al., 2010; Azevedo et al., 2008; Campbell et al., 2010; Cimino et al., 2000; Dawson and List, 2009; De Rui et al., 2015; Fedders et al., 2019; Garcia et al., 2007; Girotti et al., 2019; Hastie et al., 2005; Holmgaard et al., 2017; Isselée et al., 2001; Jones et al., 2016; Karmann et al., 2018; Kocur et al.,

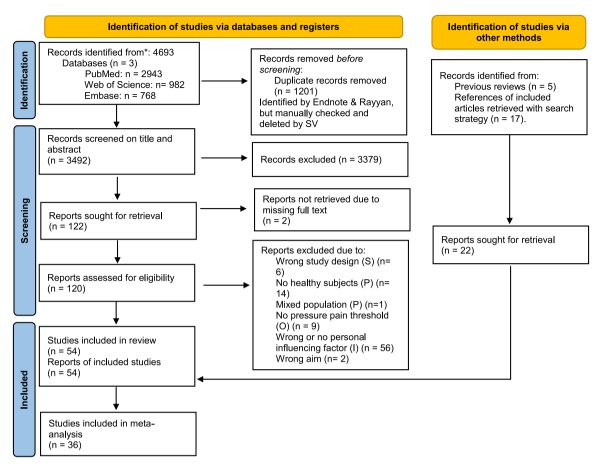


Fig. 1. PRISMA flowchart: overview of in- and exclusion process. Abbreviations: n, number.

2019; Kuppens et al., 2018; Lautenbacher et al., 2005; Lee et al., 2015; Manning and Fillingim, 2002; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Sibille et al., 2012; Teepker et al., 2010; Yang et al., 2013, 2014; You et al., 2020; Zhang et al., 2013; Shah and Luximon, 2021; Ozasa et al., 2022), and additionally, hand search vielded 22 more eligible studies (Bajaj et al., 2001; Cole et al., 2010; Edwards and Fillingim, 2001; Otto and Dougher, 1985; Rao et al., 2022; Amodei and Nelson-Gray, 1989; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Kröner-Herwig et al., 2012; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; McKendall and Haier, 1983; Petersen et al., 1992; Petrini et al., 2015; Price et al., 2013; da Silva et al., 2014; Tashani et al., 2017; Vatine et al., 1993; Donat et al., 2005; Martínez-Jauand et al., 2013). Finally, 54 studies (24 non-randomised controlled cohort (Andrzejewski et al., 2010; Dawson and List, 2009; De Rui et al., 2015; Girotti et al., 2019; Holmgaard et al., 2017; Manning and Fillingim, 2002; Pickering et al., 2002; Yang et al., 2013; Shah and Luximon, 2021; Cole et al., 2010; Edwards and Fillingim, 2001; Otto and Dougher, 1985; Rao et al., 2022; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; Petrini et al., 2015; da Silva et al., 2014; Vatine et al., 1993; Donat et al., 2005), 14 case-control (Azevedo et al., 2008; Campbell et al., 2010; Fedders et al., 2019; Kocur et al., 2019; Kuppens et al., 2018; Lautenbacher et al., 2005; Lee et al., 2015; Yang et al., 2014; You et al., 2020; Ozasa et al., 2022; McKendall and Haier, 1983; Price et al., 2013; Tashani et al., 2017; Martínez-Jauand et al., 2013), nine prospective cohort (Alves et al., 2017; Cimino et al., 2000; Garcia et al., 2007; Isselée et al., 2001; Karmann et al., 2018; Teepker et al., 2010; Zhang et al., 2013; Bajaj et al., 2001; Amodei and Nelson-Gray, 1989), and seven cross-sectional studies (Alfieri et al., 2017); Hastie et al., 2005; Jones et al., 2016; Moore et al., 2013; Shiro et al., 2017; Sibille et al., 2012;

Kröner-Herwig et al., 2012) were eligible for inclusion in this review. In case of comparative studies only data of healthy groups were used. Of these 54 studies, 36 studies were eligible for a meta-analytic approach.

Main reasons for excluding studies were the inclusion of the wrong population (no separate analysis for the healthy group) or the use of an experimentally induced setting and as such no study on personal influencing factors. Although addressing all our recourses and contacting the authors, two studies (Granges and Littlejohn, 1993; Kerem et al., 2002) were further excluded, because no full text could be found. For the screening on title and abstract, there was an agreement of 96.1% (or 3363 studies) between both reviewers, of which the remaining conflicts (3.9% or 138 studies) were resolved by consensus. An agreement of 83.2% (or 99 studies) was found between both reviewers after screening on full text. The remaining conflicts (16.8% or 21 studies) were solved by consensus (60% or 12 studies) and by contacting the last author (40% or 9 studies). Study characteristics are presented in Table 4.

3.2. Risk of bias assessment

Five (Alfieri et al., 2017; Campbell et al., 2010; Fedders et al., 2019; Girotti et al., 2019; Yang et al., 2013), nine (Azevedo et al., 2008; Dawson and List, 2009; Jones et al., 2016; Lautenbacher et al., 2005; Shiro et al., 2017; Yang et al., 2014; Petrini et al., 2015; Price et al., 2013; Tashani et al., 2017), and 40 studies (Alves et al., 2017; Andrzejewski et al., 2010; Cimino et al., 2000; De Rui et al., 2015; Garcia et al., 2007; Hastie et al., 2005; Holmgaard et al., 2017; Isselée et al., 2001; Karmann et al., 2018; Kocur et al., 2019; Kuppens et al., 2018; Lee et al., 2015; Manning and Fillingim, 2002; Moore et al., 2013; Pickering et al., 2002; Sibille et al., 2012; Teepker et al., 2010; You et al., 2020; Zhang et al., 2013; Shah and Luximon, 2021; Ozasa et al., 2022; Bajaj et al., 2001; Cole et al., 2010; Edwards and Fillingim, 2001; Otto and Dougher,

Table 4	4
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Author, year and origin	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
(Alfieri et al., 2017) Brazil	Cross-sectional	$\begin{split} N &= 75 \\ 66.8 \ y \ (4.6) \\ \varrho &= 75 \ (100\%) \end{split}$	 - 60y-75y - Female gender - Absence of psychiatric disorders - Body pain < 4 in VAS - No chronic use of analgesic or anti- inflammatory drugs - Not participating in physical therapy sessions 	Not reported	D: algometer (J Tech) S: 1 kg/s RS: pain or discomfort A: not reported P: dorsal/ventral decubitus, or sitting (depending on PPT location)	 Bilateral insertion and midbelly of following muscles: Biceps brachii Flexor capri ulnaris/ radialis Vastus medialis/lateralis Gluteus maximus 	 Functional capacity (6MWT) Handgrip strength (dynamometer) Lower limb strength (CS) 	No correlation between all factors and PPT (p $>$ 0.05).
(Alves et al., 2017) Brazil	Prospective cohort	N = 39 28.38 y (7.88) Q = 39 (100%)	 Women 18y-47y (fertile age) Healthy Normal arterial blood pressure Regular menstrual cycles (25–30 days) 	 -Irregular menstrual cycles (<25 or >30 days) Women at menopause Use of hormonal contraceptive in last 6 m Use of opioids, antidepressants or ansiolitics in last 6 m Pregnancy or attempt of pregnancy in last 6 m Breastfeeding CV, neuroendocrine, psychiatric or gynaecologic diseases Chronic or acute pain in last 6 m Central or peripheral neuropathy Physical exercises 1 h before evaluations Smoking 	D: electronic pressure algometer (Somedic) S: 50 kPa/s RS: pain A: not reported P: sitting	 Right body side: Maxillary branch of the trigeminal nerve area Forearm region 	- Menstrual cycle	PPT decreased during the phases o the menstrual cycle (p < 0.001).
(Amodei and Nelson-Gray, 1989) North Carolina	Prospective cohort	N = 12 18.42 y (SD not reported) Q = 12 (100%)	- Menstruating for at least 2 years - Menstrual cycle 20–40 days	 - Menstrual discomfort secondary to gynecological disorder of organic origin - Oral contraceptive use < 2 m prior to study participation 	D: strain-gauge pain stimulator S: not reported (standard weight 225 g, in s) RS: first pain A: not reported P: sitting	Dominant side: • Middle phalanx of index finger	- Menstrual cycle	No effect on PPT (p > 0.05).
(Andrzejewski et al., 2010) Poland	Non-randomized controlled cohort	$\begin{split} N &= 76 \\ (concerning age \\ influence) \\ Younger age \\ group (38 or 50%) \\ 22 y (SD not \\ reported) \\ Older age group: \\ (38 or 50%) \\ 65 y (SD not \end{split}$	Not reported	Not reported	D: algometer with shut-off and reset button S: 100 g/s RS: pain or discomfort A: 3 trials P: side lying	 Bilateral insertion and latent trigger points of following muscles: Superior fibular retinaculum Peroneus longus Biceps femoris Gluteus maximus/medius Tensor fascia latae Latissimus dorsi 	- Age -Level of physical activity (self-developed questionnaire)	Younger age resulted in higher PP (p < 0.05): on both sides at the insertion of: • Peroneus longus, • Biceps femoris, • Gluteus maximus, • Adductor magnus On left side at the insertion of: • Flexor capri ulnaris/radialis, On both sides at trigger points in: (continued on next page

Table 4	(continued)
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Author, year and origin	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
		reported) $\varphi = ?$ N = 22 (concerning physical activity influence) Vigorous physical activity group (16 or 73%) Moderate physical activity group (6 or 27%) Mean age not reported $\varphi = ?$				 Erector spinae Infraspinatus Levator scapulare Pectoralis major/minor Flexor carpi ulnaris/ radialis Adductor magnus 		 Flexor carpi radialis Adductor magnus On right side at trigger points in: Peroneus longus, Infraspinatus, More physical activity resulted in higher PPT (p < 0.05): On both sides at the insertion of: Superior fibular retinaculum Peroneus longus Biceps femoris Gluteus maximus/medius Tensor fascia latae Infraspinatus Pectoralis major/minor Adductor magnus On the left side at the insertion: Erector spinae On both sides at trigger points in Superior fibular retinaculum Peroneus longus Biceps femoris Gluteus maximus/medius Tensor fascia latae Infraspinatus Pectoralis major/minor Adductor magnus On the left side at the insertion: Erector scapulare Latissimus dorsi Flexor carpi ulnaris/radialis On both sides at trigger points in Superior fibular retinaculum Peroneus longus Biceps femoris Gluteus maximus/medius Latissimus dorsi Erector spinae Infraspinatus Levator scapulare Pectoralis major/minor Flexor carpi ulnaris/radialis Adductor magnus On left side at trigger point in: Tensor fascia latae Other results were non-significar (p > 0.05)
Azevedo et al., 2008) Brazil	Case-control	N = 52 Normal scapular group (26 or 50%): 22.2 y (1.2) $\varphi = 20$ Depressed scapular group (26 or 50%): 21.8 y (1.3) $\varphi = 20$	 No history of orthopaedic, neurological or dermatological conditions on the cervical spine and upper limbs for the last 12 m No use of anti- depressive medication or analgesic and anti- inflammatory drugs for the last 5 days 	after posture assessment: -Different combination of position of the anatomic landmarks	D: electronic pressure algometer (J tech) S: 3 Newton/s RS: sensation of pressure changed to pain A: 1 trial P: sitting	M. Trapezius pars descendens dominant side	-Posture (scapular position)	A depressed scapular position resulted in lower PPT (p = 0.008

Table 4	(continued)
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Author, year and origin	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
(Bajaj et al., 2001) Denmark	Prospective cohort	$\begin{split} N &= 35 \\ \wp \; 28 \; y \; (1.9) \\ \varsigma \; 30 \; y \; (1.4) \\ \wp &= 15 \; (43\%) \end{split}$	- Healthy - Normal menstrual cycle	- Dysmenorrhoea - Oral contraceptives	D: electronic algometer (Somedic) S: 30 kPa/s RS: pain or discomfort A: 3 trials P: lying	 Abdomen (T10-T12) Lower back (S2-S4) Upper arm (C7-C8) Mid-thigh (L2) 	- Menstrual cycle - Gender	Being in the ovulatory phase resulted in lower PPT measured at the lower back as compared to the other phases ($p < 0.0002$). Women presented lower PPTs at the lower back compared with men ($p < 0.01$). Other results were non-significant ($p > 0.05$).
(Campbell et al., 2015) United Kingdom	Case-control	N = 84 34 y (14.59) 9 = 38.8%	-No pain condition or medical disorder	 Active alcohol or drug abuse problem Use of narcotics, antidepressants, anticonvulsant, or muscle relaxants 	D: algometer (Somedic) S: 30 kPa/s RS: pain A: 2 trials (both sides together) P: not reported	Bilateral:M. Trapezius pars descendens	- Depression (BDI,BSI) - Catastrophizing (PCS, SCQ)	No significant correlation (p > 0.05). No significant correlation between PCS and PPT (p > 0.05) -A correlation between PPT and SC (p < 0.01, r = -0.38) was found. -SCQ predicted PPT after controllin for gender, age, ethnicity, depression and the PCS (p < 0.01, R ² = 0.32). -SCQ was a better predictor than PC (p < 0.05).
Chesterton et al., 2003) United Kingdom	Non-randomized controlled cohort	N = 240 Women 25 y (Jaber et al., 2018) Q = 120 (50%)	-Healthy	 Peripheral neuropathy Pain symptoms History of trauma or surgery to the dominant hand Current medication Diabetes Pregnancy 	D: pressure algometer (Salter Abbey Weighing Machines) S: 5 Newton/s RS: pain distinct from pressure or discomfort A: 2 trials P: not reported	Non-dominant m. interosseous dorsalis 1	-Gender	-Women reported lower PPT than men (p < 0.0005)
	Prospective cohort	N = 24 25.1 y (3.6) Q = 24 (100%)	-Regular menstrual cycle (28 ± 2 days)	 Reporting moderate or severe pain symptoms TMD and/or orofacial pain diagnosed according to the Research Diagnostic Criteria Intake of oral contraceptives Wearing of intrauterine contraceptive devises Consumption of NSAID or any other medication 1 m prior to participation Migraine Neurological disorders 	D: algometer (Somedic) S: 20 kPa/s RS: sensation of pressure changed to pain A: last 3 trials P: sitting	 Right body side (2 places): M. Masseter (M1 and M2) M. Temporalis pars anterior (T1 and T2) 	Menstrual cycle	Being in the preovulatory phase resulted in lower PPT at M1 compared to the other phases ($p < 0.05$), and at T1 compared to the luteal phase ($p < 0.05$). Other results were non-significan ($p > 0.05$).
	Non-randomized controlled cohort	Younger group (15 of 50%): 26 y (Blyth et al., 2019) Q = 8 (53%)	 Not reporting pre- existing pain Not taking any medication during testing Additional for older 	 Peripheral neuropathy Diabetes Stroke Hypertension Psychiatric disorders 	D: hydraulically driven device composed of a circular rubber probe S: 0.25 kg/cm ² /s RS: participants had to rate with a 20-point scale the JNP	Thumbnail	Age	Younger age resulted in higher P ($p < 0.05$).

Table 4 (continued)

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uthor, year and	Study Design	Participants			Device, speed, and analysis I of PPT Reported signal and position of patient	Location of PPT	Influencing factor (measurement method)	Results (p value + ES or correlation coefficient [r] if significant and if reported)
origin		Group composition and characteristics	Inclusion criteria	Exclusion				
		et al., 2018) ♀ = 6 (40%)			magnitudes that elicited alternating reports of no pain sensation (0/20) and faint pain (1/20). P: not reported			
Dawson and List, 2009) Sweden	Non-randomized controlled cohort	N (total) = 64 Middle Easterners (32 or 50%): φ 24.1 y (2.3) d 25.2 y (3.6) φ = 16 (50%) Swedes (32 or 50%): φ 23.3 y (3.3) d 24.8 y (3.5) φ = 16 (50%)	Inclusion Middle Easterners: - Participant + parents born in Iraq, Iran, Lebanon, Syria, or Palestine - Spoke first language at home Inclusion Swedish: - Born and raised in Sweden for the last two generations - Spoke Swedish at home	 Chronic pain Trauma and/or surgery to the hands Cardiac disease Analgesics or other medication that would influence pain perception Pregnancy 	D: algometer S: 30 kPa/s RS: pressure that felt like pain A: not reported P: not reported	Not reported	-Gender -Ethnicity	No difference in PPT between genders ($p > 0.05$) after pooling for cultures. No difference in PPT between Middle Easterners and Swedes ($p > 0.05$) after pooling for gende
De Rui et al., 2015) Italy	Non-randomized controlled cohort	$\begin{split} N &= 97 \\ \wp \ 71.7 \ y \ (0.6) \\ \eth \ 73.1 \ y \ (0.9) \\ \wp &= 63 \ (65\%) \end{split}$	-> 65 y -Independence in all activities of daily living, with possible exception of continence -A normal cognitive performance or mild cognitive impairment(>5/10 on SPMSQ)	 Nursing home residents Taking NSAID drugs/ analgesics in previous month Chronically treated with antidepressants or membrane-stabilizing drugs Having type 1 diabetes, systematic rheumatic diseases, tension-type headache, odontogenic pain, TMD, myopathies or fibromyalgia 	D: algometer (Fischer) S: 100 g/s RS: pain A: not reported P: not reported	 Bilateral: M. Temporalis (pars anterior, medialis and posterior) M. Masseter M. Sternocleidomastoideus M. Occipitalis M. Splenius Capitis M. Hypothenar 	-Gender -Age -Educational level (years of education)	Women reported lower PPT compared to men for all muscles $(p < 0.05)$. No difference in PPT $(p > 0.05)$. No difference in PPT $(p > 0.05)$.
Donat et al., 2005) (91) Turkey	Non-randomized controlled cohort	$\begin{split} & \text{N (total)} = 128 \\ & 60-69 \text{ y age group} \\ & (23 \text{ or } 18\%): \\ & \varphi = 16 (70\%) \\ & 70-79 \text{ y age group} \\ & (62 \text{ or } 48\%): \\ & \varphi = 38 (61\%) \\ & > 80 \text{ y age group} \\ & (43 \text{ or } 34\%): \\ & \varphi = 26 (60\%) \end{split}$	Not reported	-History of recent upper extremity injury -Any kind of orthopaedic, neurologic, or systemic pathology causing function deficiency or pain	D: dolorimeter (Wagner) S: 1 kg/s RS: pain or discomfort A: mean of 3 trials P: not reported	Bilateral: • Second finger • Fifth finger • Hand	-Age	No significant results (p > 0.05)
Edwards and Fillingim, 2001) Alabama	Non-randomized controlled cohort	N (total) $= 68$	Not reported	Not explicitly reported, but participants were screened for: -Ongoing pain problems -Hypertension -Circulatory disorders -Cardiac problems -Metabolic disease -Other significant health		Left body side:M. Trapezius pars descendensM. Masseter	-Age -CV activity (measuring of SBP and MAP in resting state)	No significant differences between age groups ($p > 0.05$) PPT correlated with, measured at r Masseter: 1. SBP ($p = 0.002$, $r = 0.50$) 2. MAP ($p = 0.005$, $r = 0.46$) In younger participants 1. SBP ($p = 0.04$, $r = 0.35$)

Author, year and	Study Design			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or	
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
				risks -Use of centrally acting agents				In older participants PPT correlated with, measured at m Trapezius: 1. SBP ($p = 0.03$, $r = 0.37$) 2. MAP ($p = 0.04$, $r = 0.37$) In younger participants 1. SBP ($p = 0.03$, $r = 0.37$) In older participants Other results were non-significant ($p > 0.05$).
(Fedders et al., 2019) Denmark	Case-control	N (total) = 40 Normal BMI group (20 or 50%): 23.30 y (1.81) $\varphi = 10$ (50%) High BMI group (20 or 50%): 27.05 y (8.53) $\varphi = 10$ (50%)	 Healthy men and women of 18–65 y Non-smokers Caucasian BMI≥ 18.5 kg/cm² 	 Pregnant or breastfeeding Pacemaker Full-grown board Lacked the ability to cooperate Performed extreme athletic activities Neurological, CV, MSK, or psychological illness Craniofacial pain, migraine, chronic TTH, new persistent headache or TMD. Dermatological skin conditions, wounds, scars or skin sensation alteration in facial region Chronic pain < 3 m or acute pain on day of study Flu or fever < 2w Addicted to drugs Consummation of alcohol 	D: handheld pressure algometer (Somedic) S: 30 kPa/s RS: sensation of pressure changed to pain A: 1 trial before CPT P: not reported	Bilateral: • M. Temporalis • M. Masseter • M. Deltoideus	-BMI (weight with SilverCrest Diagnostic Scale, Height with non- elastic measuring tape) -Gender	(p > 0.05). No significant difference in PPT between the BMI groups or gender (irrespective of BMI) (p > 0.05).
				< 24 h before study - Use of medication with impact on immune system or pain for the last 24 h - Use supplements or medication that affect body weight				
(Garcia et al., 2007) Spain	Prospective cohort	N = 30 40.94 y (14.42) Q = 18	-Not suffering from chronic or acute pain just before or during the study	 Chronic MSK disorder Frequent pain Analgesic medication in a non-sporadic manner Traumatic injury Surgery or suffered pain 6 m Psychological disorder Medical incident < 3 days before measurement (taking analgesic medication, apparition of 	D: algometer (DEP) S: 1 kg/s RS: pressure that felt like pain A: mean of 3 trials P: not reported Patients were measured in three sessions (baseline, 15 min post, and 7 days post)	Bilateral tender points of following muscles: • Trapezius pars descendens • Supraspinatus • Gluteus pars anterior Bilateral tender points of: • Occiput • Low cervical • Greater trochanter • Medial fat pad knee • Second rib • Lateral epicondyle	Gender	Woman showed lower mean PPT values than men measured at 'tota of tender points at session 3 ($p < 0.05$, ES: 0.84). Only in the first trial of session 3, woman showed lower PPT than me measured at 'total' of tender points ($p < 0.05$, ES: 0.84) Women showed lower PPT than me for the 'total' of control points ($p < 0.05$, ES: 0.85) Woman showed lower mean PPT values than men measured at control

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Author, year and Study Design Participants Device, speed, and analysis Location of PPT Influencing factor Results (p value + ES or origin of PPT Reported signal and (measurement method) correlation coefficient [r] if Inclusion criteria Exclusion Group position of patient significant and if reported) composition and characteristics medical complications or Bilateral control points: points at session 1 (p < 0.05, ES: invasive techniques) Hypothenar eminence 0.91), 2 (p < 0.05, ES: 0.74) and 3 Medial tibia (p < 0.01, ES: 0.81). Medial ulna Other results are non-significant (p > 0.05)(Girotti et al., Non-randomized N (total) = 355Not reported - MSK, skin or joint disease D: algometer (Fisher) Bilateral: -Age -PPT significantly decreased in both 2019) Carpal joint -Gender controlled cohort Younger group at site of PPT testing S: not reported (Newton/ men and women from younger age Italy (195 or 55%): - Presence of any pain cm^2) Older group to older age (p < 0.0001). 20-45 y (mean syndrome or RS: pressure became painful -BMI -Adjustment of BMI: negative A: mean of 3 trials and SD not symptomatology at the -Manual work association between age and PPT in time of assessment (difference) or mean of last 2 -Educational level (years men and woman (p < 0.0001). reported) Q = 120 (62%) - Use of pain killers trials (regression) of education) -Women had lower PPT than men in Middle aged - Severe cognitive P: not reported -Depression (GDS-sf) all age groups, but the difference group (75 or impairments or -Cognitive impairment was more attenuated with advancing 21%): neurological diseases that (SPMQS) age (young group: p = 0.001, might affect the ability to -Complex functions in middle-aged group: p = 0.02, older 46-64 y (mean discriminate between and which fully self-sufficient group: p = 0.003). and SD not reported) define painful stimuli (IADL) Gender and age influenced PPT ♀ = 35 (47%) -Comorbidities (CIRS-CI) (p = 0.03)Older group (85 or -Poly-pharmacy No difference in PPT (p > 0.05). Non-manual work resulted in lower 24%): 65–95 y (mean PPT in men (p = 0.03)and SD not No difference in PPT (p > 0.05). No difference in PPT (p > 0.05). reported) Cognitive impairment resulted in Q = 51 (60%)lower PPT in men (p = 0.04)No difference in PPT (p > 0.05). Higher CIRS-CI resulted in lower PPT in men (p = 0.02). No difference in PPT (p > 0.05). (Hastie et al., Cross-sectional N = 188-Healthy participants - Chronic pain, systemic D: handheld pressure Right body side: -Affective status No association with PPT (p > 0.05). Association between the KRS and 2012) 24.14 y (5.47) medical condition algometer (Pain Diagnostics • M. Trapezius pars (PANAS) USA Q = 110 (59%)- Use of prescription and Therapeutics) descendens -Hyper-vigilance (KRS) PPT (p < 0.01, r = 0.254). medication M. Masseter -Pain coping (CSQ) Association between the CSQ-S: 1 kg/s RS: Not reported subscale active factor and PPT Ulna A: mean of 3 trials (p < 0.05, r = -0.154).P: not reported D: handheld electronic (Holmgaard et al., Non-randomized N = 60 - No history of - Red hair • M. Tibialis anterior -Phenotypic features for No difference in PPT (p > 0.05) 2017) controlled cohort 22 y (2.30) persistent pain - History of pressure algometer M. Temporalis dark/light eve and hair A gender difference for PPT Denmark Q = ?- Phenotypic features neurophysiological or (SENSEbox, Somedic) -Gender measured at m. Temporalis for dark eye and dark mental illnesses S: 25 kPa/s (p = 0.015).RS: Pain Other results were non-significant hair colour or light eye - Ongoing pain treatment and light hair colour. - Drug or alcohol abuse A: 1 trial (before CPT) (p > 0.05)- Use of nicotine (<30days) P: not reported - Chronic headache (>2x week) - Pregnancy or nursing (Isselée et al., N = 30-Menstrual cycle All data together: PPT was different Prospective Not reported - History of medical D: algometer (Somedic Type Bilateral: 2001) cohort Group taking oral problems (diabetes, III) • M. Temporalis -Gender between gender-hormonal phases of Belgium contraceptives (10 neurological or metabolic) S: 40 kPa/s M. Masseter -Taking oral all measurements in all muscles. or 33%): - Dysmenorrhea or RS: sensation of pressure • Thumb contraceptives except for the first measurement of 26 y (SD not gynecological problems changed to pain the m. Masseter.

Table 4 (continued)

Table 4 (continued)

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Author, year and	Study Design	sign Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value $+$ ES or
rigin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
		characteristics reported) $\varphi = 10 (100\%)$ Group without contraceptives (10 or 33%): 29 y (SD not reported) $\varphi = 10 (100\%)$ Men (10 or 33%): 31 y (SD not reported) $\varphi = 0 (0\%)$		- Cycle length < 26days or > 33days	A: not reported P: not reported -Women were assessed according to their menstrual cycle during 4 sessions [2 sessions in mid-to-late follicular phase, and two in the mid-to-late follicular phase] and 10 consecutive cycles -Men were assessed weekly for 2 months and every 2 weeks thereafter for 10 months.			For women without contraceptive PPT measured at following muscle and location: • Temporalis was different between the menstru and follicular phases ($p = 0.0001$, p = 0.0009, and $p = 0.0198measurement 1,2 and 3,respectively), and between themenstrual an luteal phases(p = 0.0256), p = 0.0124 formeasurement 1 and 2, respectivel• thumbwas different between the menstruand the follicular phase (p = 0.017,measurement 1)For women taking oralcontraceptives PPT measured atfollowing locations:• M. Masseter (p = 0.0001,p = 0.0014$, and $p = 0.0038$ for measurement 2,3 and 4, respectively) • M; Temporalis ($p = 0.0001$, p = 0.0001, and $p = 0.0011$ for measurement 2,3 and 4, respectively) • thumb ($p = 0.0008$, measurement 2) was different between the menstrue phase and follicular phase compare to men. • M. Masseter ($p = 0.0001$, p = 0.0014, and $p = 0.0033$, measurement 2,3, and 4, respectively) • M. Temporalis ($p = 0.0001$, p = 0.0014, and $p = 0.0101$, for measurement 2,3, and 4, respectively) • M. Temporalis ($p = 0.0001$, p = 0.0001, and $p = 0.0101$, for measurement 2,3, and 4, respectively) • thumb (measurement 2) was different between the menstrue and ulteal phase compared to mets No difference in PPT between women and men ($p > 0.055$). No difference in PPT between
nes et al., 2016) Australia	Cross-sectional	N = 53 ç 24.4 y (3.6)	- Apparently healthy with no history of chronic pain or	Not reported	D: handheld algometer (Wagner) S: 1 kg/s	Right body side: • M. Trapezius • M. Biceps brachii	-Peak aerobic capacity (VO2peak) -Gender	women not taking medication ($p > 0.05$). Association between lower body PI and VO2peak in men ($p = 0.03$, r = -0.58). Ischemic pain tolerand

Table 4	(continued)
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	Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if
				position of patient			significant and if reported)
	Q = 17 (32%)	chronic disease - 18y-50y - Absence of a current diagnosis of depression		RS: sensation of pressure changed to pain A: mean of 3 trials P: not reported	 M. Rectus femoris M. Tibialis anterior 	-Ischemic pain tolerance (ischemic tourniquet test)	and VO2peak were predictors for this association ($p = 0.01$ and p = 0.045, respectively). No association with PPT ($p > 0.05$ No association with PPT ($p > 0.05$ Other results were non-significant ($p > 0.05$).
Prospective cohort	N = 40 38.8 y (13.5) Q = 20 (50%)	Not reported	 Acute and chronic pain Psychological disorders or medical diseases (e.g. sleep disorders) Taking psychotropic drugs or analgesics 	D: computer controlled pressure algometer (Noxitest Biomedical) S: 50 kPa/s RS: slightly painful A: mean of 5 trials P: not reported	Left body side:Fingertip of the middle fingerFingertip of the index finger	-Overnight changes (1st vs. 2nd night) -Total sleep time, efficiency, and latency -N and period of awakenings -Durations of non-REM stage & slow-wave, and REM sleep (polysomnography) -Subjective sleep quality (morning and evening DGSM)	None of the factors predicted PPT (p > 0.05).
Case-control	$\begin{split} N &= 50 \\ Normal head \\ posture group (25 \\ or 50%): \\ 39.6 y (8.1) \\ \varphi &= 25 (83\%) \\ Forward head \\ posture group (25 \\ or 50\%): \\ 40.7 y (6.8) \\ \varphi &= 25 (83\%) \end{split}$	time as average or low - 24–55 y - BMI 18.5–30 kg/m ² - VAS < 5 - NDI< 14 - Weekly working time in a sitting position of at least 35 h - Lack of orthopedic and neurological	values of anthropometric characteristics - After surgery within the thoracic and cervical spine, and, shoulder girdle	D: algometer (Wagner) S: not reported (Newton/ cm ²) RS: pain complaint A: not reported P: not reported	 M. Trapezius pars descendens M. Splenius Capitis M. Sternocleidomatoideus 	Forward head posture (photometric method)	No effect on PPT (p > 0.05).
Non-randomized controlled cohort	$\begin{split} N &= 32 \\ \wp \; 25.3 \; (5.5) \\ \eth \; 25.4 \; (4.6) \\ \wp &= 16 \; (50\%) \end{split}$	-Caucasian	 Pain in head or neck region (jaw dysfunction and headaches, subjective pain or soreness of the masticatory muscles) Currently taking medication or received other treatment General health problems or periodontal disease History of drug abuse Recent facial or cervical trauma Being in menstrual phase Smoker 	D: pressure algometer (Somedic) S: 30 kPa/s RS: pain A: 3 trials P: not reported	Bilateral: • M. Masseter • Thenar	-Age -BMI (measurement method not reported) -Gender	No effect on PPT ($p = 0.648$) No effect on PPT ($p = 0.665$) Women reported lower PPT on all locations ($p < 0.001$).
	cohort Case-control Non-randomized	cohort $38.8 \text{ y} (13.5)$ $\varphi = 20 (50\%)$ Case-control N = 50 Normal head posture group (25 or 50%): 39.6 y (8.1) $\varphi = 25 (83\%)$ Forward head posture group (25 or 50%): 40.7 y (6.8) $\varphi = 25 (83\%)$ $\varphi = 25 (83\%)$ Non-randomized N = 32 controlled cohort $\varphi = 25.3 (5.5)$ $\sigma = 25.4 (4.6)$	cohort $38.8 \text{ y} (13.5)$ $\wp = 20 (50\%)$ - Healthy individuals without any acute or chronic headaches or neck pain < 6 m $39.6 \text{ y} (8.1)$ $Q = 25 (83\%)$ - Level of physical $Q = 25 (83\%)$ - Level of physical activity in their leisure Forward head time as average or low posture group (25 or 50%): $Q = 25 (83\%)$ $Q = 25 (83\%)$ - MMI 18.5–30 kg/m² $40.7 \text{ y} (6.8)$ $Q = 25 (83\%)$ $VAS < 5$ $\varphi = 25 (83\%)$ - NDI < 14 $Weekly working timein a sitting position ofat least 35 h- Lack of orthopedicand neurologicalcomorbiditiesNon-randomizedcontrolled cohortN = 32\varphi = 25.3 (5.5)g = 25.4 (4.6)$	cohort $38.8 \ y (13.5)$ $Q = 20 (50\%)$ - Psychological disorders or medical diseases (e.g. sleep disorders) - Taking psychotropic drugs or analgesicsCase-controlN = 50 Normal head posture group (25 or 50%): $Q = 25 (83\%)$ - Healthy individuals without any acute or chronic headaches or neck pain < 6 m - Level of physical activity in their leisure forward head posture group (25 or 50%): $Q = 25 (83\%)$ - Participants with extreme values of anthropometric characteristics - After surgery within the thoracic and cervical spine, activity in their leisure im as average or low posture group (25 or 50%): $Q = 25 (83\%)$ - NDL - Level of physical activity in their leisure im a sitting position of at least 35 h - Lack of orthopedic and neurological comorbidities- Pain in head or neck region (jaw dysfunction and headaches, subjective pain or soreness of the maticatory muscles) - Currently taking medication or received other treatment - General health problems or periodontal disease - History of drug abuse - Recent facial or cervical trauma	cohort38.8 y (13.5) $Q = 20 (50\%)$ - Healthy individuals without any acute or posture group (25- Participants with extreme values of anthropometric characteristics characteristicsD: algometer (Wagner) S: 50 kPa/s $Q = 25 (83\%)$ - Healthy individuals without any acute or posture group (25- Participants with extreme threat posture group (25D: algometer (Wagner) S: not reported $Q = 25 (83\%)$ - Healthy individuals without any acute or posture group (25- Healthy individuals without any acute or posture group (25- Participants with extreme thread actes or activity in their lesior at lesion and neurological comorbiditiesD: algometer (Wagner) S: not reportedNon-randomized controlled cohortN = 32 Q = 25 (83\%)- Wack y working time in a sitting position of at least 35 h - Lack of orthopedic and neurological comorbidities- Pain in head or neck region (Jaw dysfunction and headaches, subjective pain or soreness of the masticatory muscles)D: pressure algometer (S : 30 kPa/s S: 30 kPa/sNon-randomized controlled cohortN = 32 Q = 16 (50\%)- Caucasian - Pain in head or neck region (Jaw dysfunction and headaches, subjective pain or soreness of the masticatory muscles)D: pressure algometer (S : 30 kPa/sNon-randomized controlled cohortN = 32 Q = 16 (50\%)- Caucasian - Rein in head or neck region (Jaw dysfunction and headaches, subjective A : 3 tr	cohort38.8 y (13.5) Q = 20 (50%)- Participants with extreme medical dissorders or naming psychotropic disorders) - Taking psychotropic disorders)Dressure algometer (Wagner) S: slightly painful A: mean of 5 trails P: not reported- Hengittp of the middle fingerCase-controlN = 50 posture group (25 2 = 25 (8%) a Ctivity in their leizur posture group (25 2 = 25 (8%) 4 (27, Y (6.8)- Healthy individuals and, shoulder girdle region Participants with extreme characteristics - After surgery within the region.D: algometer (Wagner) S: not reported (Newton/ P: not reported- M. Trapezius pars descendensNormal head posture group (25 2 = 25 (8%) q = 16 (50%)- Healthy individuals without any acute or posture group (25 2 = 25 (8%)- Participants with extreme characteristics - Lexel of physical and, shoulder girdle posture group (25 2 = 25 (8%)- Normal head posture group (25 2 = 25 (8%)- M. Stapella P. Not: 14 - Weekly working time in a sitting postitor or and headdens, subjective S: 30 kPa/s S: 30 kPa/s S: 30 kPa/sD: pressure algometer (Somedic) S: 30 kPa/s S: 30 kPa/sBilateral: - M. MasseterNom-candomized controlled color of a 25 4 (46) q = 16 (50%)N = 32 - Caucasian- Pain in head or neck region (law dysfunction and headdaches, subjec	cohort 38.8 y (13.5) 9 = 20 (50%) -Psychological disorders or gene diacid disoset (2, 2, 2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,

Table 4	(continued)
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Author, year and origin	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or correlation coefficient [r] if
		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	significant and if reported)
(Komiyama et al., 2007) Belgium Japan	Non-randomized controlled cohort	N = 88 Japanese (44 or 50%): \Diamond 25.0 (3.6) δ 24.7 (3.6) \Diamond = 22 (50%) Belgium (44 or 50%): \Diamond 23.9 (3.6) δ 24.4 (3.1) \Diamond = 22 (50%)	- Belgian Caucasian subjects or Japanese subjects	 Pain in head or neck region (jaw dysfunction and headaches, subjective pain or soreness of the masticatory muscles) Currently taking medication or received other treatment General health problems or periodontal disease History of drug abuse Recent facial or cervical trauma Being in menstrual phase Smoker 	D: pressure algometer (Somedic) S: 30 kPa/s RS: pain A: 3 trials P: not reported	Bilateral: • M. Masseter • Thenar	-Ethnicity -Age -Gender -Height (measurement method not reported) -Weight (measurement method not reported)	No effect on PPT ($p > 0.05$). No effect on PPT ($p > 0.05$). Women reported lower PPT on all locations ($p = 0.002$) No effect on PPT ($p > 0.05$). No effect on PPT ($p > 0.05$).
(Kröner-Herwig et al., 2012) Germany	Cross-sectional	$\begin{split} N &= 74 \\ 23.1 \ y \ (2.5) \\ \varphi &= 35 \ (47\%) \end{split}$	-Speaking German language	 Acute pain at the time of testing and/or persistent pain Consummation of alcohol on the day of the experiment Pain medication < 48 h before testing Pregnancy Bruises or injuries at the site of the pain application 	S: 1 kg/s RS: pain	Inner forearm	-Gender -Gender role (masculinity/femininity BSRI) -Pain catastrophizing (PCS) -Fear of Pain (FBQ-III and PASS) -Depression (BDI) -SBP (tourniquet) -Menstrual phase	-Women reported lower PPT than men (p = 0.001) -Gender correlated with PPT (p ≤ 0.05 , r = -0.39) No effect on PPT (p > 0.05) No correlation with PPT (p > 0.05)
(Kuppens et al., 2018) Belgium	Case-control	$\begin{split} N &= 25 \\ 26.12 \ y \ (8.29) \\ \varrho &= 13 \ (52\%) \end{split}$	 Pain-free and healthy subjects No history of shoulder or neck pain requiring medical treatment 		D: digital algometer (Wagner) S: 1 kg/s RS: pain A: not reported P: not reported	 Bilateral: M. Trapezius pars descendens dorsal side of middle finger Right body side: m. Gastrocnemius 	 Pain Vigilance and Awareness (PVAQ) Pain catastrophizing (PCS) Sports participation (participation in sport whether or not) 	No correlation with PPT for all factors ($p > 0.05$).
(Lautenbacher et al., 2005) Germany	Case-control	$\begin{split} N &= 40 \\ Younger group \\ (20 \text{ or } 50\%): \\ 27.1 \text{ y } (3.5) \\ \varphi &= 10 \ (50\%) \\ Older group (20 \text{ or } 50\%): \\ 71.6 \text{ y } (5.9) \\ \varphi &= 10 \ (50\%) \end{split}$	-Pain-free	 Conditions that could affect pain perception and report such as diabetes, hypertension, peripheral and central neuropathy, neuropsychological and psychiatric disorders. Analgesic or sedative medication < 48 h prior to test session 	D: computer-controlled pneumatic pressure algometer S + RS: steps of 200 kPa until participant reported first pain or discomfort, then decreased in steps of 100 kPa until 'no pain' was reported, followed by steps of 50 kPa until 3 upward turning points were reached. A: median value of 3 upward turning points P: not reported	Bilateral: • Middle finger • Ring finger	-Age -Gender	Younger age resulted in higher PPT (p = 0.003). No effect on PPT (p > 0.05).
(Lee et al., 1994) Korea	Non-randomized controlled cohort		 Free of pain in head and neck region at the time of the interview No tenderness to 	 Difficulty and/or pain opening mouth Sometimes stuck or locked jaw Noises in jaw joints 	S: 40 kPa/s RS: pain	Following muscles:Temporalis pars anterior, medius and posteriorMasseter pars inferior, superior and profunda	-Age -Gender	Younger age resulted in lower PPT women measured at following muscles: • Temporalis pars anterior, mediu and posterior

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Table 4 (continued)

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Author, year and	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
			digital palpation at any site	 Pain in face, cheeks, jaws, throat or temples Uncomfortable or unusual feeling of bite Frequent headache, neck and/or shoulder pain Previously treated for jaw joint problem Jaw symptoms or headache in the morning 	A: 2 trials P: not reported	 Pterygoideus medialis Digastricus posterior Sternocleidomastoideus pars medialis and superficialis Splenius capitis Trapezius insertion Trapezius pars descendens 		 Masseter pars profunda Pterygoideus medialis Digastricus posterior Sternocleidomastoideus pars superficialis Splenius capitis Trapezius insertion Trapezius pars descendens Younger age resulted in lower PPT men measured at following muscle Pterygoideus medialis Digastricus posterior Splenius capitis Trapezius insertion Trapezius pars descendens Women in their 20ies reported low PPT than men in their 20ies measured at following muscles: Temporalis pars anterior, media and posterior Masseter pars inferior, superior and profunda Digastricus posterior Sternocleidomastoideus pars medialis and superficialis Splenius capitus Trapezius insertion Trapezius pars descendens Women in their 30ies reported low PPT than men in their 30ies measured at following muscles: Masseter pars inferior, superior and profunda Digastricus posterior Sternocleidomastoideus pars medialis and superficialis Splenius capitus Trapezius pars descendens Women in their 30ies reported low PPT than men in their 30ies measured at following muscles: Masseter pars inferior, superior and profunda Pterygoideus medialis Digastricus posterior Sternocleidomastoideus pars medialis and superficialis Splenius capitis Trapezius insertion Women in their 40ies reported low PPT than men in their 40ies measured at following muscles: Masseter pars inferior, superior and profunda Pterygoideus medialis Digastricus posterior Women in their 50ies reported low PT than men in their 50ies measured at following muscles: M. masseter pars inferior, superi

Author, year and	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
(Lee et al., 2015) Taiwan	Case-control	N (total)= 20 Depressed scapular group (8 or 40%): 21.7 y (0.7) $\varphi = 3$ (38%) Normal scapular group (12 or 60%): 22.2 y (0.4) $\varphi = 7$ (58%)	Not reported	 History of orthopedic, neurological, or dermatological conditions of the cervical spine History of orthopedic conditions of upper limbs Antidepressants or any analgesics < 3 m Anti-inflammatory drugs 5days Body-implanted electronic devices 	D: electronic pressure algometer (Wagner) S: not reported (kgf/cm ²) RS: sensation of pressure changed to pain A: mean of 3 trials P: sitting	Bilateral:M. Trapezius pars descendensM. Trapezius pars transversa	Scapular position (posture assessment)	 Sternocleidomastoideus pars medialis and superficialis Depressed scapular position resulted in a lower PPT compared to normal scapular position on all locations (right m. Upper trapezius: p = 0.021, left m. Upper trapezius: p = 0.021, left m. Middle trapezius: p = 0.030, and left m. Middle trapezius: p = 0.027).
(Lemming et al., 2015) Denmark	Non-randomized controlled cohortches	N = 98 Highly active men (22 or 22%): 30.6 y (1.9) $\varphi = 0$ (0%) Normally active men (26 or 27%): 36 y (2.4) $\varphi = 0$ (0%) Highly active women (27 or 28%): 34.8 y (1.8) $\varphi = 27$ (100%) Normally active women (23 or 23%): 35.7 y (2.5)	- 20–65 y - Pain-free	 Pain in m. Trapezii Current or previous presence of pain (through brief medical history) 	D: manual pressure algometer (Somedic) S: 30 kPa/s RS: pain A: 1 trial P: not reported	Dominant side: • M. Tibialis anterior	-Gender -Physical activity level (GLTEQ)	Women reported lower PPT than men ($p = 0.019$) Highly active participants had higher PPT than normally active participants ($p = 0.049$)
(Manning and Fillingim, 2002) USA	Non-randomized controlled cohort	$\ensuremath{\wp} = 23 \ (100\%)$ N = 24 20y-24y (no info about mean age and SD) $\ensuremath{\wp} = 12 \ (50\%)$	Athlete group: - Soccer, gymnastics, football, tennis, fencing, crew, and wrestling sports Non-athlete group: -Not participating athletics on an intercollegiate level, but being physically active and participating in \geq 3 h exercise per week.	Not reported	D: pressure algometer S: 1 kg/s RS: sensation of pressure changed to pain A: mean of all trials (Mäntyselkä et al., 2001;Blyth et al., 2019;Yam et al., 2018; Backonja et al., 2009)P: not reported	Following muscles:Biceps brachiiPectoralisQuadricepsDeltoideus	-Athletic status (competition team or not) -Gender -Locus of control (RIELCS) -Ability to dissociate (DES) -Gender role attitudes (BSRI) -Coping strategies (CSQ) -Mood states (POMS and STAI) -Physical activity (questionnaire) -Number of injury experience	No overall effect of athletic status on PPT ($p > 0.05$). No overall effect of gender on PPT ($p > 0.05$), but non-athletic women had lower PPT at the m. Deltoid ($p = 0.013$), m. Pectoralis ($p = 0.028$) than non-athletic men. (ES: $0.23-0.37$) No effect on PPT ($p > 0.05$). No effect on PPT ($p > 0.05$).

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Author, year and	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
							-Pain-related self- efficacy (own design)	
Martínez-Jauand et al. (2013) Argentina	Case-control	$\begin{split} N &= 32 \\ 57.0 \ y \ (1.1) \\ \varphi &= 32 \ (100\%) \\ Early \ menopause \\ (17 \ or \ 53\%) \\ Late \ menopause \\ (15 \ or \ 47\%) \end{split}$	-Postmenopausal women -Last menstrual period > 1 y	-Pregnancy -Neurologic disease	D: digital dynamometer (Wagner) S: not reported RS: pain A: not reported P: not reported	Not reported	Menopause	No difference in PPT between earl and late menopause (p > 0.05)
Matos et al., 2011) Denmark	Non-randomized controlled cohort	$\begin{split} N &= 30 \\ 1825 \text{ y (no info} \\ about mean age \\ and SD) \\ \varrho &= 15 \ (50\%) \end{split}$	- Healthy	- Signs or symptoms of pain, hyperalgesia or allodynia in head, neck, and face region (jaw dysfunction and headaches, subjective pain or soreness of the masticatory muscles)	D: pressure algometer (Somedic) S: 30 kPa/s RS: pain A: mean of 3 trials P: not reported	Bilateral:Infraorbital foramenMental foramen	-Gender	Women reported lower PPT than men ($p = 0.006$).
McKendall and Haier, 1983) Netherlands	Case-control	$\begin{split} N &= 60 \\ \wp \; 43.78 \; y \; (11.22) \\ \eth \; 42.92 \; y \; (14.67) \\ \wp &= 40 \; (67\%) \end{split}$	Not reported	- < 20 y or > 67 y	D: pressure device with constant and equal pressure of 3 pounds S: not reported (in pounds/s) RS: time to first pain A: mean of 2 trials P: not reported	Dominant side: • First joint of index finger	-Obesity (< or > than 130% of IBW)	The obese group reached sooner th PPT compared to the non-obese group ($p < 0.01$). If mid-weight participants were excluded (obese >180% of IBW, ar non-obese <125% of IBW), obese women reported lower PPT than non-obese woman ($p < 0.05$). Other results were non-significant ($p > 0.05$).
(Moore et al., 2013) UK	Cross-sectional	$\begin{split} N &= 189 \\ 23.65 \text{ y (6.15)} \\ \varrho &= 119 \text{ (63\%)} \end{split}$	Not explicitly reported, but: Patients reported that they were not currently in pain, had no existing chronic pain condition, and were not taking analgesic medication	Not reported	D: pressure pain algometer (Somedic) S: 50 kPa/s RS: first pain A: mean of 5 trials P: not reported	Dominant side: • Back of the hand	-Gender -Pain-related fear and anxiety (PASS) -Anxiety-related symptoms (ASI-3) -Fear of different causes of pain (FPQ-III) -Pain catastrophizing (PCS) -Depression, anxiety, and stress experiences (DASS-21)	Women had lower PPT than men $(p < 0.01)$ (t-test), gender predict PPT $(p < 0.001)$, $R^2 = 0.07$) No effect on PPT $(p > 0.05)$. No effect on PPT $(p > 0.05)$.
	Non-randomized controlled cohort		Not reported	Not reported	D: focal pain stimulator S: not reported (gradually increase 640 gm) RS: participant reported a 5 on a 7-point Likert scale (slight pain) A: not reported P: not reported	Second phalanx of finger	-Gender -Masculinity /femininity (>50 BSRI)	Women reported lower PPT ($p < 0.05$). Masculinity-Femininity influenced PPT significantly ($p < 0.01$) A significant interaction between masculinity-femininity scores and gender for PPT ($p < 0.05$).
Ozasa et al., 2022) China	Case-control	N = 42 Premenopausal (21 or 50%) 45.2 y (2.4)	-Healthy	Not reported	D: not reported S: not reported RS: not reported A: not reported	Tip of the tongue	Menopause	None of the results were significat $(p > 0.05)$

Table 4 (continued)

Author, year and	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
(Petersen et al., 1992) Denmark	Non-randomized controlled cohort	postmenopausal (10 or 24%) 55.6 (2.8) Late postmenopausal (11 or 26%) 64.9 (10.8) 9 = 42 (100%) N = 40 24.2 y (4.1) 9 = 20 (50%)	- Headache free - Not taken analgesic or psychotropic drugs < 24 h before the test	Not reported	D: pressure algometer (Somedic) S: 1.1 Newton/s RS: lowest pressure that gives a sensation of pain A: 2 trials P: supine	 Bilateral: M. Temporalis with myofascial tissue M. Temporalis without myofascial tissue Middle phalanx of second finger 	-Gender -Hand dominance	Women reported lower PPT than men at all locations: • Temporalis with myofascial tisse ($p = 0.09$) • Temporalis without myofascial tissue ($p = 0.03$) • Second finger ($p < 0.05$) PPT was higher on the right finger compared to the left finger in righ handed participants ($p = 0.07$). Other results were non-significant ($p > 0.05$).
(Petrini et al., 2015) Denmark	Non-randomized controlled cohort	$\begin{split} N &= 40 \\ Younger group \\ (20 \text{ or } 50\%): \\ 24.6 \text{ y } (3.6) \\ \varphi \ 10 \ (50\%) \\ Older group \ (20 \text{ or } 50\%): \\ 73.6 \text{ y } (6.6) \\ \varphi &= 10 \ (50\%) \end{split}$	- Cognitive intact capabilities (MMSE score 28–30) - Pain-free	 Conditions that could affect pain perception and pain report Presence of severe ongoing pain Neuropsychological and psychiatric disorders Diabetes Signs of rheumatic or arthritic disease especially on 	D: electronic hand-held pressure algometer (Somedic) S: 30 kPa/s RS: pain A: mean of 4 trials P: not reported	Bilateral: • Index finger • M. Trapezius	-Age -Gender	Elderly men reported lower PPT compared to young men ($p < 0.001$). Young women reported lower PPT compared to young men ($p < 0.001$). Other results were non-significant ($p > 0.05$)
				hand/fingers and neck/ shoulders -Taken any analgesic or sedative < 48 h prior to test				
(Pickering et al., 2002) France	Non-randomized controlled cohort	N = 42 Younger group: 9 22 y (Raja et al., 2020) $\delta 22 y$ (Mäntyselkä et al., 2001) Older group: 9 74 y (Blyth et al., 2019) $\delta 74 y$ (Yam et al., 2018) Total of $9 = 21$ (50%)	Inclusion: - 18y-25y (young group) or > 70 y (older group) - Living in the community - Physically and psychologically autonomous -No overt history of cardiovascular Disease - No medication, especially no analgesics	 Any disorder likely to affect sensory function, neurological and psychiatric antecedents Cutaneous illness of upper limbs Alcohol and street drug consumption 	D: electronic pressure algometer (Somedic) S: 1.1 Newton/s RS: pain A: mean of 4 trials P: not reported	 Second finger Third finger Fourth finger Fifth finger 	-Age -Gender -Cognitive capacities (MMSE) -Psychomotor performance (CRT)	Younger age in men resulted in higher PPT in men ($p < 0.001$). No effect on PPT ($p > 0.05$). No effect on PPT ($p > 0.05$). Correlation between PPT and CRT ($p = 0.01$, $r = -0.52$)

Table 4	(continued)
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Author, year and	Study Design	Participants			Device, speed, and analysis of PPT Reported signal and	Location of PPT	Influencing factor	Results (p value + ES or correlation coefficient [r] if significant and if reported)
origin		Group composition and characteristics	Inclusion criteria	Exclusion	position of patient		(measurement method)	
Price et al., 2013) Canada	Case-control	$\begin{split} N &= 40 \\ Obese group (20 \\ or 50\%): \\ 26.3 (7.1) \\ \varphi &= 10 (50\%) \\ Non-obese group \\ (20 or 50\%): \\ 26.5 y (7.4) \\ \varphi &= 10 (50\%) \end{split}$	of tobacco, tea or coffee Extra inclusion (older group): - No decreased finger mobility - No sign of rheumatic or arthritic disease - No pain or uncomfortable sensation on passive and active hand and finger movements - Obesity (BMI ≥ 30 kg/cm ²) (obese group) - Non-obesity (control group)	 < 18 y or > 45 y Habitual use of recreational drugs, tobacco or alcohol Regular or frequent night shift work Presence or history of psychiatric or neurological disorder Diabetes Hypertension Chronic pain Any other severe medical condition as these conditions may be associated with altered pain sensitivity 	RS: pain	Non-dominant side: • Thenar eminence • Thenar thumbnail	-Obesity (measurement method not reported)	No effect on PPT (p > 0.1).
Rao et al., 2022) Bhopal	Non-randomized controlled cohort	$\begin{split} N &= 95 \\ & \text{Group 1 (25 or} \\ & 26\%): \\ & 18y-21y (no info \\ & about mean age \\ & and SD) \\ & \varphi &= 25 (100\%) \\ & \text{Group 2 (20 or} \\ & 21\%): \\ & 30y-40y (no info \\ & about mean age \\ & and SD) \\ & \varphi &= 20 (100\%) \\ & \text{Group 3 (25 or} \\ & 26\%) \\ & 25y-41y (no info \\ & about mean age \\ & and SD) \\ & \varphi &= 25 (100\%) \\ & \varphi &= 25 (100\%) \\ & \text{Group 4 (25 or} \\ & 26\%) \\ \end{split}$	- Physically and mentally healthy Group 1: studying medical laboratory technology Group 2: housewives and working women Group 3: family planning center and taking oral contraceptives Group 4: medical students	- Consumption of analgesic, antidepressant, anxiolytic drug or hormones	D: metallic aerated water bottle cap and a sphygmomanometer S: 4 mm/Hg RS: pricking pain A: mean of 3 trials (group 1–3), mean of 9 trials (group 4) P: not reported	Flexor surface forearm	Menstrual cycle	PPT varied related to age and menstrual cycle in women. PPT was higher in phase I and menstruation in women taking contraceptives, while PPT was only at phase II of the menstrua cycle in women taking no oral contraceptives (p < 0.001).

Table 4 (cor	ntinued)
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Author, year and	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or	
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)	
Shah and Luximon (2021) China	Non-randomized controlled cohort	about mean age and SD) $\varphi = 0 (0\%)$ N = 218 φ 27.79 y (11.55) d 26.36 y (9.24) $\varphi = 109 (50\%)$	Healthy	-Facial soft tissue or bone deformities	D: a hand-held ultrasound indentation device S: less than 2 mm/s RS: pain A: mean of 3 trials P: not reported	 23 landmarks on head- and forehead region 13 landmarks on frontal part of the face 16 landmarks on both sides of the face 8 landmarks on the nek region 	-Age -Gender -BMI	Age correlated with 27 landmarks of the face region in women ($p < 0.05$) Women reported lower PPT in 67 out of 76 landmarks compared to men ($p < 0.05$). BMI correlated with 12 landmarks of the face region in men ($p < 0.05$). Other results were non-significant ($p > 0.05$).	
(Shiro et al., 2017) Japan	Cross-sectional	$\begin{split} N &= 86 \\ 20.9 \ y \ (0.8) \\ \varrho &= 43 \ (50\%) \end{split}$	- 20y-29y - No ongoing pain problems	-History of chronic pain conditions and serious health conditions such as neurological diseases or diabetes -Use of sedatives, analgesics, or other medications.	D: mechanical pressure algometer (Digital Force Gage) S: 5 Newton/s RS: pain because of pressure A: not reported P: sitting	Dominant side:M. Extensor carpi radialis brevis	-Gender -BMI (scale) -BMR (formula) -Moderate-to-vigorous physical activity (accelerometer past 7 days) transformed to METs	Women had lower PPT than men (p = 0.019). No effect on PPT (p > 0.05). BMR was a predictor of PPT in the overall study population (p = 0.018, r^2 =0.054), but not if accounted for gender (p > 0.05). No effect on PPT (p > 0.05).	
(Sibille et al., 2012) Florida	Cross-sectional	$\begin{split} N &= 372 \\ 23.7 \text{ y (SD not} \\ reported) \\ \varrho &= 205 \end{split}$	 - 18y-45y - No report of clinical pain - No psychiatric disturbance or substance disorder - No use of tobacco products or centrally acting medications 	Not reported	D: handheld algometer (Pain Diagnostics and Therapeutics) S: 1 kg/s RS: pain because of pressure A: mean of 3 trials P: semi-lying	 Right body side: M. Trapezius pars descendens M. Masseter pars superficialis ulna 	Affect balance style (PANAS)	No significant results (p > 0.05).	
(da Silva et al., 2014) Brazil	Non-randomized controlled cohort	N = 126 49.4 (23.7) Q = 65 (52%)	- Healthy	 Previous trauma or surgery to the face or skull Orofacial and/or generalized pain Neuropathic conditions Neurodegenerative diseases Neuroendocrine or rheumatological diseases Neural infections Chronic use of medication Nasal obstructions Allergies or other upper respiratory, gustative abnormalities Issues before the tests that might interfere with the results 	D: electronic pressure algometer (Somedic) S: 50 kPa/s RS: pain A: not reported P: sitting	Bilateral: • Front • Cheek • Chin • Anterior tibia skin	-Age -Gender	Participants between 61 and 75 y reported higher PPT compared to participants between 18 and 30 y and between 45 and 60 y (p < 0.001). Women reported higher PPT compared to men (p < 0.001).	
(Tashani et al., 2017) UK	Case-control	N = 74 Normal BMI (25 or 33%): 28.3 y (9.3)	Not reported	with the results -Pre-existing medical condition - Currently seeking medical care	D: handheld algometer (Somedic) S: 10 kPa/s RS: pain	Non-dominant side: • Thenar eminence	BMI (scale)	The obese group had lower PPT compared to the overweight group (p = 0.005) and the normal range group $(p = 0.001)$.	

Table 4	(continued	1)
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	Study Design	Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or	
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)	
		Overweight BMI (24 or 33%): 32.7 y (9.1) Obese BMI (25 or 33%: 36.3 (7.5) Total of $\varphi = 37$ (50%)		 Taking medication Experienced pain < 6 m Previously diagnosed with a chronic pain condition Experiencing disturbances in skin sensation Dermatological condition Pregnant Regularly undertake vigorous exercise 	A: mean of last 3 trials P: not reported			Obese and normal BMI women reported lower PPT compared to overweight women ($p = 0.002$). Higher BMI predicted lower PPT ($p = 0.019$, $r^2 = 0.074$) Other results were non-significant ($p > 0.05$).	
Feepker et al., 2010) Germany	Prospective cohort	$\begin{split} N &= 32 \\ 27.3 \ y \ (6.1) \\ \varphi &= 32 \ (100\%) \end{split}$	 Regular menstrual cycle (28 ± 1 day) Examined by a neurologist and psychologist Not taking any drugs or oral contraceptives on a regular basis No use of analgesics and sedatives < 24 h before testing 	 Pregnancy Hypertension Acute and chronic pain Endocrine, gynecological, or psychiatric disorders Peripheral and central neuropathy Dermatosis at the test side 	D: pressure algometer (Somedic) S: 10 kPa/s RS: slight pain A: mean of last 3 trials P: not reported	Left body side: • Forearm	Menstrual cycle	PPT was higher on the 22nd day compared to the 1st day of menstrual cycle ($p = 0.004$, Bonferroni: $p = 0.015$) Other results were non-significant ($p > 0.05$).	
/atine et al., 1993) Israël	Non-randomized controlled cohort		-Right-hand dominant	-Systemic disease or pain syndrome	D: pressure algometer S: 1 kg/s RS: pain A: not reported P: not reported	Bilateral:Mastoid processesExternal malleolusSternum (2 locations)	Gender	No significant differences at all locations ($p > 0.05$).	
Yang et al., 2013) Denmark China		$\begin{split} N &= 58 \\ \text{Danish group (29} \\ \text{or 50\%):} \\ 27.0 \text{ y (5.0)} \\ \varphi &= 15 (50\%) \\ \text{Chinese group (29} \\ \text{or 50\%):} \\ 28.2 \text{ y (4.0)} \\ \varphi &= 15 (50\%) \end{split}$	 No experience with similar tests Born and raised in their home country without migration ≥ 3 y of university education 	-Ongoing pain or reports of chronic pain < 6 m - Serious systemic diseases - Previous radiotherapy or chemotherapy - Intake of medicine affecting the central nervous system - Fibromyalgia - Self-reported bruxism or psychogenic illnesses	•	Bilateral: • M. Masseter • Mandibula	-Ethnicity -Gender	Chinese participants showed lowe total (both PPT locations summed PPT than Danes ($p < 0.001$, ES: 0.227). Women showed lower total PPT the men ($p < 0.001$, ES 0.184)	
Yang et al., 2014) Denmark	Case-control	N = 70 42.3 y (12.5) Q = 36 (51%)	-Healthy	 Participated in any kind of clinical test similar to the present one Ongoing pain or chronic pain < 6 m Systemic diseases or previous radio or chemotherapy Taken any medicine Iweek that affects the nervous system Physical or mental disorders 	D: computerized pressure algometer S: 30 kPa/s RS: first pain A: not reported P: not reported	Bilateral: • M. Masseter • M. Temporalis • Thenar	-Age -Gender	None of the results were significan (p > 0.05).	

Table 4 (con	ntinued)
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Author, year and	Study Design	Participants				Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
(You et al., 2020) USA	Case-control	$\begin{split} N &= 177 \\ Group 1 (43 \text{ or } 24\%): \\ 18.7 \text{ y } (1.2) \\ \varphi &= 23 (53\%) \\ Group 2 (23 \text{ or } 13\%): \\ 20.3 \text{ y } (2.0) \\ \varphi &= 6 (26\%) \\ Group 3 (50 \text{ or } 28\%): \\ 20.1 \text{ y } (2.5) \\ \varphi &= 25 (50\%) \\ Group 4 (25 \text{ or } 14\%): \\ 19.6 \text{ y } (2.3) \\ \varphi &= 11 (44\%) \\ Group 5 (36 \text{ or } 20\%): \\ 19.8 \text{ y } (1.8) \\ \end{split}$	-Healthy adults 18y- 30y Group 1: abstainers Group 2: moderate drinkers with recent drinkers with recent drinkers without recent drinking episode Group 4: binge drinkers with a recent drinking episode Group 5: binge drinkers without recent drinking	 Women: being in their menstrual period Any chronic physical and mental health issues Prescription medication use (except contraceptives and vitamins) Needle phobia Injury or skin condition on the feet (pain testing site) 	D: handheld algometer (Wagner) S: Not explicitly reported (according to Rolke et al.: 30 kPa/s) RS: not reported A: mean of 3 trials P: Not reported	Non-dominant side: • Dorsum foot	-Drinking history -Standard drinks and hours spent drinking for a typical week (DDQ)	PPT was different between group ($p < 0.001$). Group 5 had lower PPT compared group 3 ($p = 0.013$). Moderate drinking was associated with an increase, binge drinking was associated with a decrease in PPT Group 4 showed reduced PPT compared to all the other groups ($p < 0.005$). Other results were non-significant ($p > 0.05$).
(Zhang et al., 2013) China	Epidemiological cohort	$\varphi = 21 (58\%)$ N = 2517 18y-85y (No info about mean age and SD) $\varphi = 0 (0\%)$	-Men	 History of rheumatoid arthritis, gout, diabetes, or cancer History of chronic prostatitis Anti-inflammation medication < 4weeks 	D: digital pressure algometer (Wagner) S: 1 kg/cm2/s RS: unpleasant pain sensation A: mean of both sides (1 trial) P: not reported	Bilateral: • M. Triceps • Inguinal line	-Age -BMI (scale and stadiometer) -Waist circumference -SBP and DBP (sphygmomanometer) -Education (interview) -Manual occupation (interview) -Leisure time physical exercise (interview) -Smoking (interview) -Drinking (interview) -Obesity within age groups	Younger age resulted in lower PF (p < 0.001) on both locations. -Lower BMI resulted in higher PF (p < 0.001) at m. Triceps. -Correlation between BMI and PF at m. Triceps (p < 0.001, r = -0.119) -Smaller waist circumference resulted in higher PPT on both locations (p < 0.001) -Correlation between waist circumference and PPT at m. Trice (p < 0.001, r = -0.150). -The PPT at m. Triceps had a negative association with waist -Lower PPT at m. Triceps was for in men with waist circumference \geq 90 cm in all age groups (p < 0. (95%CI= -0.122 to -0.038). -A SBP (p = 0.005) and DPB (p = 0.048) difference between participants with a lower, middle bicher PT at m. and the lower, middle

higher PPT at inguinal line. -Correlation between DBP and PPT at m. Triceps (p = 0.041, r = -0.041).

-Higher level of education resulted in lower PPT on both locations $(p < 0.001) \label{eq:prod}$

Author, year and	Study Design	Design Participants			Device, speed, and analysis	Location of PPT	Influencing factor	Results (p value + ES or
origin		Group composition and characteristics	Inclusion criteria	Exclusion	of PPT Reported signal and position of patient		(measurement method)	correlation coefficient [r] if significant and if reported)
								- Higher level of education resulted in lower PPT on both locations (p < 0.001) after age and BMI adjustment. -Non-manual occupation resulted i lower PPT on both locations (p < 0.001), also after age and BM adjustment (p < 0.001). -Moderate or high leisure time physical exercise resulted in lower PPT (p < 0.001 for m. Triceps, p = 0.006 for Inguinal line), also after age and BMI adjustment on both locations (p < 0.001). No effect on PPT (p > 0.05). -Drinking more alcohol resulted in higher PPT at m. Triceps (p < 0.000 or age and BMI adjusted: p = 0.008). -Central obesity resulted in lower PPT on both locations (p < 0.001) and inguinal line (p = 0.008). -Central obesity resulted in lower PPT on both locations (p < 0.001) and m. Triceps, p = 0.001 at inguinal line) after age and BMI adjustment -Lower PPT at m. Triceps in total obesity men only in age group ≥ 40 y.

Abbreviations: 6MWT, 6-minute walk-test; A, analysis; ASI-3, Anxiety Sensitivity Index; BDI, Beck Depression Index; BMI, body mass index; BMI, Body Mass Index; BMR, Basal Metabolic Rate; BSI, Brief Symptom Inventory; BSRI, Bem Gender Role Inventory; CIRS-CI, Cumulative Illness Rating Scale - Comorbidity Index; cm, centimetre; CPT, Cold Pressure Test; CRT, choice reaction time; CS, 30 s Chair-to-stand-test; CSQ, Coping Strategies Questionnaire; CV, cardiovascular; D, device; DASS-21, Depression Anxiety Stress Questionnaire-21; DBP, diastolic blood pressure; DDQ, Daily Drinking Questionnaire; DES, Dissociative Experiences Scale; DGSM, German Sleep Society; DGSM, German Sleep Society; ES, effect size; FPQ-III, Fear of Pain Questionnaire; FPT, filament pin-prick pain threshold; g, gram; GDS-sf, Geriatric Depression Scale-short form; GLTEQ, Godin Leisure-Time Exercise Questionnaire; h, hour; IADL, Instrumental Activities of Daily Living; IBW, Ideal Body Weight; JNP, Just Noticeable Pain; kg, kilogram; kPa, kilopascal; KRS, Kohn Reactivity Scale; m., muscle; MAP, mean arterial blood pressure; METs, Metabolic Equivalents; MMSE, Mini-mental test; MSK, musculoskeletal; N, Newton; N, sample size; NDI, Neck Disability Index; NSAID, Nonsteroidal anti-inflammatory agents; P, position; PANAS, Positive and Negative Affect Scale; PANAS, Positive and Negative Affect Scale; PASS, Pain Anxiety Symptoms Scale; PCS, Pain Catastrophizing Scale; PMS, Profile of Moods States; PPT, pressure pain threshold; PPT, Pressure Pain Threshold; PVAO, Pain Vigilance and Awareness Questionnaire; REM, rapid-eve movement; RIELCS, Rotter Internal-External Locus of Control Scale; RS, reported signal; s, second; S, speed (increase in force); SBP, systolic blood pressure; SBP, systolic blood pressure; SCQ, Situational Catastrophizing Questionnaire; SD, Standard Deviation; SES, Schmerzempfindungsskala; SPMSQ, Short Portable Mental Status Questionnaire; STAI, State Trait Anxiety Inventory; TDT, tactile detection threshold; TMD, Temporomandibular Disorder; TTH, tension type headache; VAS, Visual Analogue Scale; y, years of age

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1985; Rao et al., 2022; Amodei and Nelson-Gray, 1989; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Kröner-Herwig et al., 2012; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; McKendall and Haier, 1983; Petersen et al., 1992; da Silva et al., 2014; Vatine et al., 1993; Donat et al., 2005; Martínez-Jauand et al., 2013) received a low, moderate and high RoB score, respectively. After scoring, reviewers agreed on 57.1% of the domains (Kappa value: 0.34) and 67.8% of the sub-domains (Kappa value: 0.41), but a full agreement was achieved after discussion. In most studies the domain "Study attrition" and "Study confounding" were not applicable due to the study design, and therefore not considered for the judgement of overall RoB score. Limited reporting of study participation in cross-sectional and

Table 5

Quality assessment.

case-control studies were the main reasons for attaining a higher RoB score (Table 5). Regarding the LoE, four studies received a LoE 3 (Dawson and List, 2009; Girotti et al., 2019; Yang et al., 2013; Petrini et al., 2015), **39 studies a LoE 4** (Alfieri et al., 2017; Alves et al., 2017; Andrzejewski et al., 2010; Azevedo et al., 2008; Campbell et al., 2010; Cimino et al., 2000; De Rui et al., 2015; Fedders et al., 2019; Garcia et al., 2007; Holmgaard et al., 2017; Isselée et al., 2001; Jones et al., 2016; Karmann et al., 2018; Lautenbacher et al., 2005; Manning and Fillingim, 2002; Pickering et al., 2002; Shiro et al., 2017; Teepker et al., 2010; Yang et al., 2014; Zhang et al., 2013; Shah and Luximon, 2021; Bajaj et al., 2001; Cole et al., 2010; Edwards and Fillingim, 2001; Otto and Dougher, 1985; Rao et al., 2022; Amodei and Nelson-Gray, 1989; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al.,

Study	1	2	3	4	5	6	Overall RoB	LoE
(Alfieri et al., 2017)	Low	N/A	Low	Moderate	N/A	Low	Low	4
(Alves et al., 2017)	High	High	Low	Moderate	High	Low	High	4
(Amodei and Nelson-Gray, 1989)	High	High	Moderate	Moderate	N/A	Moderate	High	4
(Andrzejewski et al., 2010)	High	N/A	Moderate	Moderate	N/A	Moderate	High	4
(Azevedo et al., 2008)	Moderate	N/A	Low	Moderate	N/A	Low	Moderate	4
(Bajaj et al., 2001)	High	High	Moderate	Moderate	N/A	High	High	4
(Campbell et al., 2015)	Moderate	N/A	Low	Low	N/A	Low	Low	4
(Chesterton et al., 2003)	High	N/A	Low	Moderate	N/A	Low	High	4
(Cimino et al., 2000)	High	High	Moderate	Low	High	Moderate	High	4
(Cole et al., 2010)	High	N/A	Low	Moderate	N/A	Moderate	High	4
(Dawson and List, 2009)	Low	N/A	Low	Moderate	N/A	Moderate	Moderate	3
(De Rui et al., 2015)	Moderate	N/A	Low	Moderate	N/A	Moderate	High	4
(Donat et al., 2005)	Moderate	N/A	Low	Moderate	N/A	Moderate	High	4
(Edwards and Fillingim, 2001)	High	N/A	Low	Moderate	N/A	Moderate	High	4
(Fedders et al., 2019)	Low	N/A	Low	Moderate	N/A	Low	Low	4
(Garcia et al., 2007)	High	High	Moderate	Low	High	Low	High	4
(Girotti et al., 2019)	Moderate	N/A	Low	Low	N/A	Low	Low	3
(Hastie et al., 2012)	Moderate	N/A	Moderate	Moderate	N/A	Low	High	5
(Holmgaard et al., 2017)	High	N/A	Low	Moderate	N/A	Moderate	High	4
(Isselée et al., 2001)	High	High	High	Moderate	High	Low	High	4
(Jones et al., 2016)	Low	N/A	Moderate	Moderate	N/A	Low	Moderate	4
(Karmann et al., 2018)	High	High	Low	Moderate	N/A N/A	Low	High	4
	-	N/A	Moderate		N/A N/A	Low	High	4 5
(Kocur et al., 2019)	High			Moderate			U	5 4
(Komiyama and De Laat, 2005)	High	N/A	Low	Moderate	N/A	Moderate	High	
(Komiyama et al., 2007)	High	N/A	Low	Moderate	N/A	Low	High	4
(Kröner-Herwig et al., 2012)	High	N/A	Low	Moderate	N/A	Low	High	5
(Kuppens et al., 2018)	High	N/A	Moderate	Moderate	N/A	Moderate	High	5
(Lautenbacher et al., 2005)	Moderate	N/A	Low	Moderate	N/A	Low	Moderate	4
(Lee et al., 1994)	High	N/A	Low	Moderate	N/A	Moderate	High	4
(Lee et al., 2015)	Moderate	N/A	Moderate	Moderate	N/A	Low	High	5
(Lemming et al., 2015)	Moderate	N/A	Moderate	Moderate	N/A	Moderate	High	4
(Manning and Fillingim, 2002)	High	N/A	Moderate	Moderate	N/A	Moderate	High	4
(Martínez-Jauand et al., 2013)	High	N/A	Moderate	Moderate	N/A	Moderate	High	5
(Matos et al., 2011)	High	N/A	Low	Moderate	N/A	Low	High	4
(McKendall and Haier, 1983)	High	N/A	Moderate	High	N/A	Moderate	High	5
(Moore et al., 2020)	High	N/A	Low	Moderate	N/A	Low	High	5
(Otto and Dougher, 1985)	High	N/A	Moderate	Moderate	N/A	Moderate	High	4
(Ozasa et al., 2022)	High	N/A	Moderate	High	N/A	Low	High	5
(Petersen et al., 1992)	High	N/A	Moderate	Moderate	N/A	Low	High	4
(Petrini et al., 2015)	Moderate	N/A	Low	Moderate	N/A	Low	Moderate	3
(Pickering et al., 2020)	High	N/A	Moderate	Moderate	N/A	Low	High	4
(Price et al., 2013)	Low	N/A	Moderate	Moderate	N/A	Low	Moderate	4
(Rao et al., 2022)	High	High	High	Moderate	High	Moderate	High	4
(Shah and Luximon, 2021)	High	N/A	Low	Moderate	N/A	Low	High	4
(Shiro et al., 2017)	Moderate	N/A	Low	Moderate	N/A	Low	Moderate	4
(Sibille et al., 2012)	High	N/A	Low	Moderate	N/A	Moderate	High	5
(da Silva et al., 2014)	High	N/A	Low	High	N/A	Low	High	4
(Tashani et al., 2017)	Moderate	N/A	Low	Moderate	N/A	Low	Moderate	4
(Teepker et al., 2010)	Moderate	Moderate	Low	Moderate	N/A	Moderate	High	4
(Vatine et al., 1993)	High	N/A	Low	Moderate	N/A	Moderate	High	4
(Yang et al., 2013)	Low	N/A N/A	Low	Moderate	N/A N/A	Low	Low	3
(Yang et al., 2013)	Low	N/A N/A	Low	Moderate	N/A N/A	Moderate	Moderate	4
(You et al., 2020)	High	N/A N/A	Moderate	High	N/A N/A	Moderate	High	4 5
	0			0			•	
(Zhang et al., 2013)	Low	High	Moderate	Moderate	N/A	Low	High	4

Abbreviations: LoE= Level of Evidence, N/A= not applicable

Bias due to 1 = Study Participation, 2 = Study attrition, 3 = Prognostic factor measurement, 4 = Outcome measurement, 5 = Study Confounding, 6 = Statistical Analysis and Reporting

2007; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; Price et al., 2013; da Silva et al., 2014; Tashani et al., 2017; Vatine et al., 1993; Donat et al., 2005), and 11 studies a LoE 5 (Hastie et al., 2005; Kocur et al., 2019; Kuppens et al., 2018; Lee et al., 2015; Moore et al., 2013; Sibille et al., 2012; You et al., 2020; Ozasa et al., 2022; Kröner-Herwig et al., 2012; McKendall and Haier, 1983; Martínez-Jauand et al., 2013). None of the prospective cohort studies scored LoE 3, because their overall RoB was high.

3.3. PPT measurement & locations

Forty-six studies (Alfieri et al., 2017; Alves et al., 2017; Andrzejewski et al., 2010; Azevedo et al., 2008; Campbell et al., 2010; Cimino et al., 2000; Dawson and List, 2009; De Rui et al., 2015; Fedders et al., 2019; Garcia et al., 2007; Girotti et al., 2019; Hastie et al., 2005; Holmgaard et al., 2017; Isselée et al., 2001; Jones et al., 2016; Karmann et al., 2018; Kocur et al., 2019; Kuppens et al., 2018; Lautenbacher et al., 2005; Lee et al., 2015; Manning and Fillingim, 2002; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Sibille et al., 2012; Teepker et al., 2010; Yang et al., 2013, 2014; You et al., 2020; Zhang et al., 2013; Bajaj et al., 2001; Edwards and Fillingim, 2001; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Kröner-Herwig et al., 2012; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; Petrini et al., 2015; Price et al., 2013; da Silva et al., 2014; Tashani et al., 2017; Vatine et al., 1993) used an algometer as device to determine PPT (Alves et al., 2017; Campbell et al., 2010; Cimino et al., 2000; Fedders et al., 2019; Holmgaard et al., 2017; Isselée et al., 2001; Moore et al., 2013; Pickering et al., 2002; Teepker et al., 2010; Bajaj et al., 2001; Edwards and Fillingim, 2001; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; Petrini et al., 2015; Tashani et al., 2017). A strain-gauge pain stimulator (Amodei and Nelson-Gray, 1989), a focal pain stimulator (Otto and Dougher, 1985), a digital dynamometer (Martínez-Jauand et al., 2013), a dolorimeter (Donat et al., 2005), a hand-held ultrasound indentation device (Shah and Luximon, 2021), a hydraulically driven device (Cole et al., 2010) and a water bottle cap and sphygmomanometer (Rao et al., 2022) were used in the other studies. Kilopascal (kPa) served as the most prevalent unit (Alves et al., 2017; Campbell et al., 2010; Cimino et al., 2000; Dawson and List, 2009; Fedders et al., 2019; Holmgaard et al., 2017; Isselée et al., 2001; Karmann et al., 2018; Lautenbacher et al., 2005; Moore et al., 2013; Teepker et al., 2010; Yang et al., 2013, 2014; You et al., 2020; Bajaj et al., 2001; Edwards and Fillingim, 2001; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petrini et al., 2015; da Silva et al., 2014; Tashani et al., 2017), followed by kilogram force (kgf) (Alfieri et al., 2017; Andrzejewski et al., 2010; De Rui et al., 2015; Garcia et al., 2007; Hastie et al., 2005; Jones et al., 2016; Kuppens et al., 2018; Lee et al., 2015; Sibille et al., 2012; Zhang et al., 2013; Cole et al., 2010; Otto and Dougher, 1985; Amodei and Nelson-Gray, 1989; Kröner-Herwig et al., 2012; Price et al., 2013; Vatine et al., 1993; Donat et al., 2005), Newton (N) (Azevedo et al., 2008; Girotti et al., 2019; Kocur et al., 2019; Pickering et al., 2002; Shiro et al., 2017; Chesterton et al., 2003; Petersen et al., 1992), millimetres of mercury (mmHg) (Rao et al., 2022), pounds (McKendall and Haier, 1983) and millimetres (mm) (Shah and Luximon, 2021). Speed of pressure administration differed between 20 kPa/s (Cimino et al., 2000) and 50 kPa/s (Alves et al., 2017; Karmann et al., 2018; Moore et al., 2013; da Silva et al., 2014), between 0.1 kg/s (Andrzejewski et al., 2010; De Rui et al., 2015) and 1 kg/s (Alfieri et al., 2017; Garcia et al., 2007; Hastie et al., 2005; Jones et al., 2016; Kuppens et al., 2018; Manning and Fillingim, 2002; Sibille et al., 2012; Zhang et al., 2013; Kröner-Herwig et al., 2012; Vatine et al., 1993; Donat et al., 2005), between 1.1 N/s (Pickering et al., 2002; Petersen et al., 1992) and 5 N/s (Shiro et al., 2017; Chesterton et al., 2003), 4 mmHg/s (Rao et al., 2022), less than 2 mm/s (Shah and Luximon, 2021), or was not given (Girotti et al., 2019; Kocur et al., 2019; Lee et al., 2015; Amodei and Nelson-Gray,

1989; McKendall and Haier, 1983; Martínez-Jauand et al., 2013). There was a wide variety in locations assessed, but locations could be pooled into different body regions: lower arm/hand region (Alfieri et al., 2017; Alves et al., 2017; Andrzejewski et al., 2010; Girotti et al., 2019; Hastie et al., 2005; Isselée et al., 2001; Karmann et al., 2018; Kuppens et al., 2018; Lautenbacher et al., 2005; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Teepker et al., 2010; Yang et al., 2014; Cole et al., 2010; Rao et al., 2022; Amodei and Nelson-Gray, 1989; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Kröner-Herwig et al., 2012; McKendall and Haier, 1983; Petersen et al., 1992; Petrini et al., 2015, 2015; Price et al., 2013; Tashani et al., 2017; Donat et al., 2005), shoulder girdle/upper arm region (Alfieri et al., 2017; Andrzejewski et al., 2010; Campbell et al., 2010; De Rui et al., 2015; Fedders et al., 2019; Jones et al., 2016; Kocur et al., 2019; Kuppens et al., 2018; Manning and Fillingim, 2002; Zhang et al., 2013; Bajaj et al., 2001; Lee et al., 1994; Petrini et al., 2015; Vatine et al., 1993), neck region (Azevedo et al., 2008; Campbell et al., 2010; De Rui et al., 2015; Hastie et al., 2005, 2005; Kocur et al., 2019; Kuppens et al., 2018; Lee et al., 2015; Sibille et al., 2012; Edwards and Fillingim, 2001; Lee et al., 1994; Petrini et al., 2015), leg region (Alfieri et al., 2017; Andrzejewski et al., 2010; Holmgaard et al., 2017; Jones et al., 2016; Kuppens et al., 2018; Manning and Fillingim, 2002; Zhang et al., 2013; Bajaj et al., 2001; Lemming et al., 2015; Vatine et al., 1993), face region (Alves et al., 2017; Cimino et al., 2000; De Rui et al., 2015; Fedders et al., 2019; Hastie et al., 2005; Holmgaard et al., 2017; Isselée et al., 2001; Yang et al., 2013, 2014; Shah and Luximon, 2021; Edwards and Fillingim, 2001; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Matos et al., 2011; Petersen et al., 1992; Vatine et al., 1993) or other (e.g. tongue (Ozasa et al., 2022), head (Shah and Luximon, 2021), or different body regions pooled by the authors of the included studies or if only one study examined a specific body region (Garcia et al., 2007); da Silva et al., 2014). Calculation and reported signal to stop the measurement was comparable between studies, but details can be found in Table 4.

3.4. (Possible) influencing factors

Activities of daily living (ADL) (Girotti et al., 2019), age (Andrzejewski et al., 2010; De Rui et al., 2015; Girotti et al., 2019; Lautenbacher et al., 2005; Pickering et al., 2002; Yang et al., 2014; Zhang et al., 2013; Shah and Luximon, 2021; Cole et al., 2010; Edwards and Fillingim, 2001; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Petrini et al., 2015; da Silva et al., 2014; Donat et al., 2005); alcohol consumption (You et al., 2020; Zhang et al., 2013), blood pressure (Zhang et al., 2013; Edwards and Fillingim, 2001; Kröner--Herwig et al., 2012), body mass index (BMI) (Fedders et al., 2019; Girotti et al., 2019; Shiro et al., 2017; Zhang et al., 2013; Shah and Luximon, 2021; Komiyama and De Laat, 2005; McKendall and Haier, 1983; Price et al., 2013; Tashani et al., 2017), cognitive factors (depression (Campbell et al., 2010; Manning and Fillingim, 2002; Moore et al., 2013; Kröner-Herwig et al., 2012), pain catastrophizing (Campbell et al., 2010; Hastie et al., 2005; Kuppens et al., 2018; Moore et al., 2013; Kröner-Herwig et al., 2012), pain vigilance (Hastie et al., 2005; Kuppens et al., 2018; Kröner-Herwig et al., 2012), affect (Hastie et al., 2005; Sibille et al., 2012), cognitive capacities (Girotti et al., 2019; Manning and Fillingim, 2002; Pickering et al., 2002), fear (Moore et al., 2013), self-efficacy (Manning and Fillingim, 2002), and ability to dissociate and locus of control (Manning and Fillingim, 2002)), comorbidity (Girotti et al., 2019; Manning and Fillingim, 2002), contraceptives (Isselée et al., 2001), hair colour (Holmgaard et al., 2017), education (De Rui et al., 2015; Girotti et al., 2019; Zhang et al., 2013), ethnicity (Dawson and List, 2009; Yang et al., 2014; Komiyama et al., 2007), forward head posture (Kocur et al., 2019), function (Alfieri et al., 2017), gender (Dawson and List, 2009; De Rui et al., 2015; Fedders et al., 2019; Garcia et al., 2007; Girotti et al., 2019; Holmgaard et al., 2017; Isselée et al., 2001; Jones et al., 2016; Lautenbacher et al., 2005; Manning and

Table 6

Strength of Recommendation table (qualitative approach).

Influencing factor	Study	Location of PPT measurement	Influence (+) or not (-)	Direction of influence	Level of evidence	Subgroup SoR (if possible)	Overall SoR
lge	Andrzejewski, 2010 (41)	Lower arm/hand	+	Younger = higher PPT	4 7		1
0-	Cole, 2010 (72)	region	+	Younger = higher PPT	4		
	Girotti, 2019 (49)		+	Younger = higher PPT	4		
	Lautenbacher, 2005 (57)		+	Younger = higher PPT	4		
	Petrini, 2015 (86)		+	Younger = higher PPT	3		
	Pickering, 2002 (61)		+	Younger = higher PPT	4	Conflicting	
	Donat, 2005 (91)		-		4		
	Komiyama, 2005 (78) Komiyama, 2007 (79)		-		4		
	Yang, 2014 (66)				4		
	Lee, 1994 (81)	Shoulder girdle/arm	+	Older = higher PPT	4 7		
	Zhang, 2013 (68)	region*	+	Older = higher PPT	4	 Conflicting 	
	Andrzejewski, 2010 (41)		+	Younger = higher PPT	4		
	Lee, 1994 (81)	Neck region	+	Older = higher PPT	4]		
	Petrini, 2015 (86)		+	Younger = higher PPT	3		Conflicting
	De Rui, 2015 (46)		-		4	 Conflicting 	
	Edwards, 2001 (73)		-		4		
	Shah, 2021 (69)	Face region	-+	Older - higher DDT	4		
	Lee, 1994 (81) Shah, 2021 (69)	Face region	+/-	Older = higher PPT Older = higher PPT (only in	4		
	Shall, 2021 (03)			women)	4		
	De Rui, 2015 (46)		-		4	Conflicting	
	Edwards, 2001 (73)		-		4	B	
	Komiyama, 2005 (78)		-		4		
	Komiyama, 2007 (79)		-		4		
	Yang, 2014 (66)		-		4	-	
	Zhang, 2013 (68)	Leg region	+	Older = higher PPT	4	Conflicting	
	Andrzejewski, 2010 (41)		+	Younger = higher PPT	4 _	J	
	Shah, 2021 (69)	Head region	-		4		
	Da Silva, 2014 (88)	Face region + Tibia	+	Older = higher PPT	4	-	
lcohol	You, 2020 (67)	Foot	+	More alcohol = lower PPT	5		Conflicting
Consumption	Zhang, 2013 (68)	Upper arm, inguinal line	+	More alcohol = higher PPT	4		J .
lood pressure*	Zhang, 2013 (68)	Shoulder girdle/upper	+	Association PPT and SBP, not DBP	4	-]
	Kröner-Herwig, 2012 (80)	arm region	_	HOLDBP	5_	 Conflicting 	
	Edwards, 2001 (73)	Neck, face region	+	Higher SBP and MAP =	4	2	Conflicting
	Luwarus, 2001 (75)	Neek, face region	'	higher PPT	-		conneting
	Zhang, 2013 (68)	Inguinal line	+	Higher SBP and DBP = higher PPT	4	-]
MI*	McKendall, 1982 (84)	Lower arm/hand	+	Lower BMI = higher PPT	5 -	-]
	Tashani, 2017 (89)	region	+	Lower BMI = higher PPT	4		
	Price, 2013 (87)	-	-	5	4	Conflicting	
	Shiro, 2013 (62)		-		4		
	Girotti, 2019 (49)		-		3		
	Komiyama, 2005 (78)		-		4]	
	Zhang, 2013 (68)	Shoulder girdle/upper	+	Lower BMI = higher PPT	4	_ Conflicting	 Conflicting
	Fedders, 2019 (47)	arm region*	-		4 _	L L	
	Shah, 2021 (69) Komiyama, 2005 (78)	Face region	+/-		4 4	- Weak SoR	
	Fedders, 2019 (47)		-		4	for no influence	
	Shah, 2021 (69)	Neck region	-		4 -	J ISI IIS IIII dence	
	Shah, 2021 (69)	Head region	-		4		
	Zhang, 2013 (68)	Inguinal line	-		4]
ognitive factor							
Ability to dissociate	Manning, 2002 (59)	Shoulder girdle/upper arm region, thigh	-		4		Preliminary Sol for no influenc
Affect	Sibille, 2012 (63)	Shoulder, neck, face,	-		5	-	Very weak SoR
	Hastie, 2005 (97)	lower arm region	-		5	-	 for no influence
						٦	
Catastrophizing	Campbell, 2010 (43)	Shoulder girdle, neck	+/-	Only CSQ predicted PPT	4		
-	Hastie, 2005 (97)	region	+	Association CSQ – PPT	5	- Conflicting	
	Kummana 2010 (EC)		_		5		
	Kuppens, 2018 (56) Hastie, 2005 (97)	Lower arm/hand		Association CSQ – PPT	5	1	

Table 6 (continued)

kröner-Hervig, 2012 (80) Moore, 2013 (60) Lower arm/hand region - 5 - Very weak SOR for no influence for no influence Fear Moore, 2013 (60) Hand - 5 - Very weak SOR for no influence Preliminary So for no influence Locus of control Manning, 2002 (59) Shoulder girdle/upper arm region, thigh - 4 - Preliminary So for no influence Pain vigilance* Hastie, 2005 (97) Shoulder, neck, face, lower arm region + Higher PPT 5 - - Conflicting Self-efficacy Girotti, 2019 (80) Shoulder girdle/upper arm region, thigh - More comorbidities = lower PPT 3 - - Conflicting Juccation Zhang, 2013 (68) Upper arm, inguinal line + Higher education = lower PPT 4 -							
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Hastie, 2005 (97) Face regin • Association CSQ - PPT 5 Cagnitive impairment Circuit, 2013 (49) Hand region • Cognitive lower PPT 5 Depression Carnificing • Conflicting • • Conflicting • Conflicting •				-			
Kuppers, 2018 (50) Lower Fig - 5 Cognitive impairment Grintl, 2019 (49) Hand region - Cognitive impairment = lower PPT 3 biowr PPT - Conflicting Conflicting Depression Campbell, 2019 (43) Shoulder gridle/ unpairment, 2002 (59) Shoulder gridle/ marring, 2002 (59) - - 4 - Conflicting Moder, 2013 (60) Hand - <th< td=""><td></td><td></td><td>Face region</td><td>+</td><td>Association CSO - PPT</td><td></td><td></td></th<>			Face region	+	Association CSO - PPT		
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Picketing, 2002 (61)		Girotti, 2019 (49)	Hand region	+	Cognitive impairment =	3]
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Depression Campbell, 2010 (43) Manning, 2002 (19) Soulder girlel (1000m -1000m (2001)) Moder at 500 (1000m (2001)) Moder at 5000m (2001) Moder at 5000m (2001)			Chardelan sindle (on a s	-		-	Conflicting
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Manning. 2002 (59) Upper arm region 4 influence influence For no Moore. 2013 (60) Warmingan (2002 (50) Usper arm region 5 Very weak Soft for no influence For no Moore, 2013 (60) Manning. 2002 (59) Shoulder girlde/upper - 4 Preliminary Soft for no influence Pain vigilance* Massing. 2002 (59) Shoulder, neck, face, influence + Higher hypervigilance = 5 - Pain vigilance* Haste, 2005 (97) Shoulder, neck, face, influence + Higher hypervigilance = 5 -	Depression	Campbell, 2010 (43)	Shoulder girdle/	-		4 🗍 Weak SoR for no	,
Moore, 2013 (60) region -	•						Moderate SoR
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Table 6 (continued)

Face region* (1) 8) 9) Head region Neck region Shoulder, neck, upper leg region, knee, olbew lewer leg rife	+ + + + + + +	lower PPT Women = lower PPT	4 4 4 4 4 4 4 4 4 4 4 4	Conflicting	
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1) 8) 9) Head region Neck region Shoulder, neck, upper leg region, knee,	+ + + + +	Women = lower PPT Women = lower PPT	4 4 4 4 4 4 4		
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Shoulder, neck, upper leg region, knee,	+ +	Women = lower PPT Women = lower PPT	4 4		
leg region, knee,	+	Women = lower PPT	4		
elbow, lower leg, ribs					
pooled Face, lower leg pooled	+	Women = higher PPT	4		
Abdomen, lower back	+/-	(only at lower back, women = lower PPT)	4		
?	-		4		7
Finger	+	Influence on PPT	4		1
Arm, shoulder, thigh 2 (80) Forearm	-	/	4 5		Conflicting
(41) Leg region	+	Higher PA = higher PPT	4	7	٦
(,	+	Higher PA= lower PPT	4		
)	+	Higher PA = higher PPT	5		
	+	Higher VO2peak = lower PPT in men	4	 Conflicting 	
	-		5		
	-		4	ļ	
(41) Shoulder girdle/upper arm region	+++	Higher PA = higher PPT Higher PA = lower PPT	4 4		- Conflicting
annregion	-	Higher FA - lower FFT	4	Conflicting	Connicting
I	-		4		
	-		5		
Lower arm/hand region	+/-	/ (only BMR predicted PPT)	4	Conflicting	
(41)	+	Higher PA = higher PPT	4	Conneting	
	-		5		
(41) Back	+	Higher PA= higher PPT	4]
013 ?	-		5		Very weak SoR
Tongue	-		5		for no influence
Face region	+	PPT decreased during the	4		7
	+	menstrual cycle PPT decreased during the monstrual cycle	4	_ Conflicting	
	+	menstrual cycle Being in preovulatory phase = lower PPT	4		See following
Lower arm/hand region	+	PPT decreased during the menstrual cycle	4	See following	page
region	+	menstrual cycle PPT decreased during the menstrual cycle	4	page	
	+	First phases of menstrual	4		ך
	+	PPT increased during the	4	- Conflicting	Conflicting
	-	/	4		
	+	Being in ovulatory phase = lower PPT	4	-	
	Abdomen, lower back, upper arm, thigh Neck, shoulder region,	+ - Abdomen, lower back, + upper arm, thigh	cycle = higher PPT + PPT increased during the menstrual cycle - / Abdomen, lower back, + Being in ovulatory phase = upper arm, thigh lower PPT	cycle = higher PPT + PPT increased during the 4 menstrual cycle - / 4 Abdomen, lower back, + Being in ovulatory phase = 4 upper arm, thigh lower PPT	cycle = higher PPT + PPT increased during the 4 Conflicting menstrual cycle - / 4 Abdomen, lower back, + Being in ovulatory phase = 4 upper arm, thigh Iower PPT

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Table 6 (continued)

	Lee, 2015 (58)	upper back Neck, shoulder region, upper back	+	lower PPT Depressed scapular = lower PPT	5	Weak SoR for influence
Hand dominance	Petersen, 1992 (85)	Face, finger	+	Right handed = higher PPT on right side	4	Preliminary SoR for influence
Psychomotor performance	Pickering, 2002 (61)	Finger	+	Better psychomotor performance = lower PPT	4	Preliminary SoR for influence
Manual work	Zhang, 2013 (68)	Upper arm, inguinal line	+	Non-manual work = lower PPT	4	Preliminary SoR for influence
ADL	Girotti, 2019 (49)	Wrist	-		3	Preliminary_SoR for no influence
Forward head posture	Kocur, 2019 (55)	Neck, shoulder region	-	/	5	Preliminary SoR for no influence
Contraceptives	Isselée, 2001 (52)	Face, thumb	-	/	4	Preliminary SoR for no influence
Hair colour	Holmgaard, 2017 (51)	Face, lower leg	-	/	4	Preliminary SoR for no influence
Functional capacity/strength	Alfieri, 2017 (39)	Upper and lower arm, Upper leg	-	/	4	Preliminary SoR for no influence
Sleep	Karmann, 2018 (54)	Finger	-	/	4	Preliminary SoR for no influence
Polypharmacy	Girotti, 2019 (49)	Wrist	-	/	3	Preliminary SoR for no influence
Smoking	Zhang, 2013 (68)	Upper arm, inguinal line	-	/	4	Preliminary SoR for no influence

Studies in **bold** could be included in the meta-analyses. * = meta-analysis is dominant over qualitative analysis.

Abbreviations: SoR, Strength of Recommendation; PPT, pressure pain threshold; ADL, activities of daily living; BMI, Body Mass Index; CSQ, Coping Strategies Questionnaire; PCS, pain catastrophizing scale; SBP, systolic blood pressure; DBP, diastolic blood pressure; MAP, mean arterial pressure; PA, physical activity

Fillingim, 2002; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Yang et al., 2013, 2014; Shah and Luximon, 2021; Bajaj et al., 2001; Otto and Dougher, 1985; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Kröner-Herwig et al., 2012; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; Petrini et al., 2015; da Silva et al., 2014; Vatine et al., 1993), gender role (the gender that one associates with, not biologically related) (Manning and Fillingim, 2002; Otto and Dougher, 1985; Kröner-Herwig et al., 2012), hand-dominance (Petersen et al., 1992), level of physical activity (Andrzejewski et al., 2010; Jones et al., 2016; Kuppens et al., 2018; Manning and Fillingim, 2002; Shiro et al., 2017; Zhang et al., 2013; Lemming et al., 2015), menopause (Ozasa et al., 2022; Martínez-Jauand et al., 2013), menstrual cycle phase (Alves et al., 2017; Cimino et al., 2000; Isselée et al., 2001; Teepker et al., 2010; Bajaj et al., 2001; Rao et al., 2022; Amodei and Nelson-Gray, 1989), polypharmacy (Girotti et al., 2019), psychomotor performance (Pickering et al., 2002), scapular position (Azevedo et al., 2008; Lee et al., 2015), sleep (Karmann et al., 2018), smoking (Zhang et al., 2013), and manual work (Zhang et al., 2013) were examined in the included studies as possible personal influencing factors for PPT.

Most factors were only examined in univariate analysis, except for some studies investigating the influence of age (controlled for gender and measurement site (Lautenbacher et al., 2005), gender and BMI (Girotti et al., 2019), and gender and site (Yang et al., 2014)), alcohol consumption (controlled for age and BMI (Zhang et al., 2013)), cognitive factors (controlled for gender, age, and/ or ethnicity (Campbell et al., 2010); Girotti et al., 2019; Moore et al., 2013; Pickering et al., 2002), comorbidity, (controlled for gender (Girotti et al., 2019)), education (controlled for gender and/ or BMI (Girotti et al., 2019); Zhang et al., 2013), ethnicity (controlled for gender (Yang et al., 2013); Komiyama et al., 2007), gender (controlled for ethnicity, site, age, measurement site and gender role (Lautenbacher et al., 2005); Yang et al., 2013, 2014; Komiyama et al., 2007; Kröner-Herwig et al., 2012), gender role (controlled for gender (Otto and Dougher, 1985); Kröner-Herwig et al., 2012), level of physical activity (controlled for gender (Lemming et al., 2015)), psychomotor performance (controlled for age and cognitive impairment (Pickering et al., 2002)), manual work and smoking (controlled for age and BMI (Zhang et al., 2013)), ADL (which controlled for gender (Girotti et al., 2019)) and sleep (controlled for measurement trial (Karmann et al., 2018)).

3.5. Meta-analytic and qualitative analysis of influencing factors

All clinically measurable personal influencing factors are summarized in a strength of recommendation table (Table 6). In addition, forest plots of the meta-analyses are added and studies were included if sufficient data was present (Figs. 2–14). Results of univariate analysis were used to make conclusions in both the meta-analytic and qualitative approach, in order to create an overview per personal factor. Only for the study of Campbell et al (Campbell et al., 2010). the multivariate analysis was used (no reporting of univariate analysis). For the meta-analytic approach, the 95% confidence interval (CI) in Zhang et al (Zhang et al., 2013). and standard error of mean in Rao et al (Rao et al., 2022). was recalculated as standard deviation according to the method described in the handbook of Cochrane (7.7.3.2 Obtaining standard deviations from standard errors, 2021).

If at least 2/3 of the studies could be included in the (subgroup) meta-analysis, the meta-analysis was dominant and the studies not

	Y	ounger			Older			Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
8.2.1 Hand region									
Girotti 2019 Carpal joint	50.45	24.65	195	27.8	13.13	160	9.7%	1.11 [0.89, 1.34]	+
Lautenbacher 2005 Finger	438.25	278.58	20	253.75	157.61	20	8.0%	0.80 [0.15, 1.45]	
Pickering 2002 Finger	517	156	21	490.5	160	21	8.2%	0.16 [-0.44, 0.77]	
Subtotal (95% CI)			236			201	26.0%	0.74 [0.16, 1.32]	◆
Heterogeneity: Tau ² = 0.20; Chi ² =	8.62, df =	2 (P = 0.	01); l² :	= 77%					
Test for overall effect: Z = 2.49 (P =	= 0.01)								
8.2.2 Shoulder/arm region									
Lee 1994 Trapezius insertion	302	61.7	128	339	75.5	39	9.3%	-0.57 [-0.93, -0.20]	
Zhang 2013 Triceps	5.69	1.87		6.27	1.85	532	10.0%	-0.31 [-0.41, -0.21]	•
Subtotal (95% CI)			2113			571	19.3%	-0.38 [-0.59, -0.16]	•
Heterogeneity: Tau ² = 0.01; Chi ² =	1.76, df =	1 (P = 0.	18); l² :	= 43%					
Test for overall effect: Z = 3.39 (P =		v	- ,, -						
8.2.3 Neck region									
Edwards 2001 Upper trapezius	662.5	97.2	34	569.36	59.9	34	8.7%	1.14 [0.63, 1.66]	
Lee 1994 Neck region	262.55	58.23		285.38	60.25	39	9.3%	-0.39 [-0.75, -0.03]	
Subtotal (95% CI)	202100	00.20	162	200100	00120	73	18.0%	0.37 [-1.13, 1.86]	
Heterogeneity: Tau² = 1.12; Chi² = : Test for overall effect: Z = 0.48 (P =		= 1 (P < (0.0000°	l); l² = 96	%				
8.2.4 Face region									
Edwards 2001 Masseter	239.6	25.5	34	196.6	19.6	34	8.4%	1.87 [1.29, 2.44]	
Lee 1994 Face region	198.17	38	128	278.63	62.06	39	9.1%	-1.79 [-2.20, -1.38]	
Subtotal (95% CI)			162			73	17.5%	0.03 [-3.55, 3.62]	
Heterogeneity: Tau² = 6.64; Chi² = Test for overall effect: Z = 0.02 (P =		= 1 (P <	0.0000)1); I² = 9	9%				
8.2.5 Other									
da Silva 2014 Face region + Tibia	307.98	109.62	57	370.02	157.16	68	9.3%	-0.45 [-0.80, -0.09]	
Zhang 2013 Inguinal line	3.3	1.35	1985	3.98	0.69	532	10.0%	-0.55 [-0.64, -0.45]	*
Subtotal (95% Cl)			2042			600	19.3%	-0.54 [-0.63, -0.45]	♦
Heterogeneity: Tau² = 0.00; Chi² = Test for overall effect: Z = 11.35 (P	,	· ·	60); l² :	= 0%					
Total (95% CI)			4715			1518	100.0%	0.06 [-0.35, 0.47]	•
Heterogeneity: Tau ² = 0.44; Chi ² =	329.60 df	= 10 (P	< 0.000	$(001) \cdot 1^2 =$	97%				
Test for overall effect: $Z = 0.30$ (P =		10 (1	0.000		5.70				-4 -2 0 2 4
est for subgroup differences: Chi ²			– ი იიი)4) I ² – 8	0.7%				Younger Older
reactor aubgroup unterences. Chi-	- 20.72, 0	u = 4 (P	- 0.000	/+/, I = 0	0.1 /0				

Fig. 2. Meta-analysis for the influence of age (mean differences + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

	No/rarely drinking Drinking							Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% Cl
You 2020 Dorsum foot	3.35	1.44	44	3.06	1.42	133	7.8%	0.29 [-0.20, 0.78]	
Zhang 2013 Inguinal line	3.45	1.35	1131	3.57	1.51	1395	49.3%	-0.12 [-0.23, -0.01]	-=-
Zhang 2013 Triceps	5.86	1.68	1131	6.07	1.83	1395	42.9%	-0.21 [-0.35, -0.07]	
Total (95% CI)			2306			2923	100.0%	-0.13 [-0.27, 0.02]	•
Heterogeneity: Tau ² = 0.01	; Chi² = 4.	09, df =	2 (P =	0.13); l²	= 51%	, 0		-	-1 -0.5 0 0.5 1
Test for overall effect: Z = 1	.72 (P = 0	0.09)							No/rarely drinking Drinking

Fig. 3. Meta-analysis for the influence of alcohol consumption (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

included in the meta-analysis are described separately. Otherwise, both the findings of the meta-analytic and the qualitative approach are described (all articles included in both approaches). If subgroup analyses per body region were possible, the result is given, otherwise only an overall result (not taken into account the body region of the PPT measurement) is presented.

Results are described following the methods described in Santesso et al. (Santesso et al., 2020).

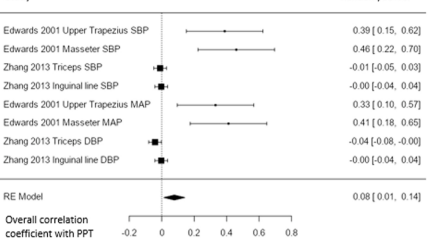
3.6. Age

Influencing factor: PPT measured at shoulder girdle/upper arm region. Three studies evaluated the influence of age on PPT measured at the shoulder/arm region, Fig. 2 shows the results of the meta-analysis of two studies (Zhang et al., 2013; Lee et al., 1994) and displays that older age may result in a slight higher PPT measured at the shoulder girdle/upper arm region compared to younger age (SMD: -0.38, 95%CI: -0.59; -0.16). A third study could not be implemented in the meta-analysis, but also reported an influence, however in the opposite direction (Andrzejewski et al., 2010) (Table 6).

Conflicting results: overall PPT and PPT measured in all body region subgroups, except shoulder girdle/upper arm region. Sixteen studies investigated the influence of age on PPTs (Andrzejewski et al., 2010; De Rui et al., 2015; Girotti et al., 2019; Lautenbacher et al., 2005; Pickering et al., 2002; Yang et al., 2014; Zhang et al., 2013; Shah and Luximon, 2021; Cole et al., 2010; Edwards and Fillingim, 2001; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Petrini et al., 2015; da Silva et al., 2014; Donat et al., 2005). Fig. 2 presents the meta-analysis of seven studies (Girotti et al., 2019; Lautenbacher et al., 2005; Pickering et al., 2002; Zhang et al., 2013; Edwards and Fillingim,

Random, 95%CI

Study



Heterogeneity: Tau²=0.0053; H²= 6.959, df= 7.000 (P < 0.001); l²= 85.63% Test for overall effect: Z= 2.34 (P= 0.019)

Fig. 4. Meta-analysis for the influence of blood pressure (correlation coefficients). Abbreviations: RE, random effects; PPT, pressure pain threshold; SBP, systolic blood pressure; MAP, mean arterial blood pressure, DBP, diastolic blood pressure.

2001; Lee et al., 1994; da Silva et al., 2014) and showed probably no overall influence of age on PPT (SMD: 0.06; 95%CI: -0.35; 0.47). However, subgroup analyses revealed that being of older age may result in having a lower PPT in the lower arm/hand region (SMD: 0.74; 95%CI: 0.16; 1.32) (Girotti et al., 2019; Lautenbacher et al., 2005; Pickering et al., 2002). More than half of the studies could not be integrated in the meta-analysis, and as such, also qualitative description including all 16 studies was necessary and showed an overall and subgroup-related conflicting strength of recommendation or conclusion (Table 6).

3.7. Alcohol consumption

Non-influencing factor. Two studies examined the influence of alcohol consumption on PPT, Fig. 3 shows the meta-analysis (You et al., 2020; Zhang et al., 2013). The frequency of alcohol consumption may have no influence on PPT (SMD: -0.13, 95%CI: -0.27; 0.02) (Fig. 3).

3.8. Blood pressure

Influencing factor. Three studies examined the effect of systolic (Zhang et al., 2013; Edwards and Fillingim, 2001; Kröner-Herwig et al., 2012), diastolic (Edwards and Fillingim, 2001) and mean arterial blood pressure (Zhang et al., 2013) on PPT, Fig. 4 shows the results of the meta-analysis, including two studies (Zhang et al., 2013; Edwards and Fillingim, 2001). Having higher blood pressure values may be associated with a slight higher PPT (CC= 0.08, 95%CI: 0.01; 0.14). The remaining study reported no influence (Kröner-Herwig et al., 2012). However, this study was of the lowest evidence (level 5) (Table 6).

3.9. BMI

Non-influencing factor. Nine studies investigated the influence of higher BMI on PPT (Fedders et al., 2019; Girotti et al., 2019; Shiro et al., 2017; Zhang et al., 2013; Shah and Luximon, 2021; Komiyama and De Laat, 2005; McKendall and Haier, 1983; Price et al., 2013; Tashani et al., 2017), of which six had sufficient data to perform a meta-analysis (Fedders et al., 2019; Zhang et al., 2013; Shah and Luximon, 2021; McKendall and Haier, 1983; Price et al., 2013; Tashani et al., 2017). Having a higher BMI probably result in no difference in PPT compared to normal BMI regarding mean differences and standard deviations

(Fedders et al., 2019; Zhang et al., 2013; McKendall and Haier, 1983; Price et al., 2013; Tashani et al., 2017) (Fig. 5, SMD: -0.20, 95%CI: -0.46; 0.05) and regarding correlation coefficients (Fedders et al., 2019; Zhang et al., 2013; Shah and Luximon, 2021) (Fig. 6, CC: -0.00, 95%CI: -0.08; 0.08) (Fig. 6). Also subgroup meta-analyses (lower arm/hand region [SMD: -0.56, 95%CI: -1.17, 0.04] and shoulder girdle/upper arm region [SMD: -0.15, 95%CI: -0.53; 0.22]) likely showed no influence. The three remaining studies (Girotti et al., 2019; Shiro et al., 2017; Komiyama and De Laat, 2005) also reported no influence at the lower arm/hand and face region (Table 6).

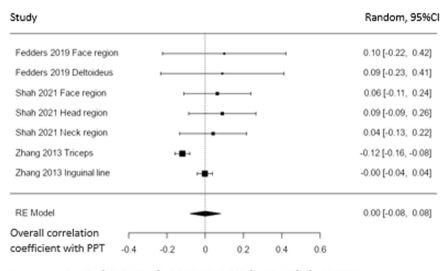
3.10. Cognitive factors

Non-influencing factor: pain vigilance, affect, depression, fear, selfefficacy, ability to dissociate and locus of control. Three studies evaluated the influence of pain vigilance on PPT, and two studies (Kröner--Herwig et al., 2012; Hastie et al., 2012) were included in the meta-analysis (Fig. 7). According to the meta-analysis, pain vigilance may not be associated with PPT (CC: 0.02, 95%CI: -0.26; 0.30), however the evidence is very uncertain. The third study also showed no influence on PPT (Kuppens et al., 2019) (Table 6). Two studies examined the influence of affect on PPT at different body regions (Hastie et al., 2005; Sibille et al., 2012). This qualitative analysis suggests no influence (Table 6). Four studies evaluated the influence of depression on PPT (Manning and Fillingim, 2002; Moore et al., 2013; Kröner-Herwig et al., 2012; Campbell et al., 2015). This analysis probably resulted in no overall influence. Also subgrouping revealed no influence (Table 6). Furthermore, the results of the qualitative analysis about fear (Moore et al., 2013), self-efficacy (Manning and Fillingim, 2002), ability to dissociate (Manning and Fillingim, 2002) and locus of control (Manning and Fillingim, 2002) showed no influence on PPT, but is preliminary, because the conclusion is based on only one study per personal factor (Table 6).

Conflicting results: pain catastrophizing and cognitive impairment. Five studies included pain catastrophizing as possible influencing factor for PPT (Campbell et al., 2010; Kröner-Herwig et al., 2012; Hastie et al., 2012; Kuppens et al., 2019; Moore et al., 2020). Meta-analysis, including three studies (Campbell et al., 2010; Kröner-Herwig et al., 2012; Hastie et al., 2012), revealed that pain catastrophizing may have no overall influence on PPT measured at different body regions (Fig. 8, CC: -0.11,

		Obese		Nor	n-obese			Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl
5.2.1 Hand region									
McKendall 1982	81.88	56.09	26	131.03	51.14	34	11.7%	-0.91 [-1.45, -0.37]	
Price 2013	3.8	1.15	20	3.7	1.15	20	10.0%	0.09 [-0.53, 0.71]	
Tashani 2017	620.72	423.81	25	1,219.92	923.04	25	10.8%	-0.82 [-1.40, -0.24]	
Subtotal (95% CI)			71			79	32.5%	-0.56 [-1.17, 0.04]	\bullet
Heterogeneity: Tau ² = 0.20;	Chi ² = 6.5	52, df = 2	(P = 0	.04); l ² = 69	9%				
Test for overall effect: Z = 1.	.83 (P = 0	.07)							
5.2.2 Arm/shoulder region									
Fedders 2019 Deltoideus	276.69	134.63	20	254.19	113.87	20	10.0%	0.18 [-0.44, 0.80]	
Zhang 2013 Triceps	5.41	1.7	357	5.9	1.86	2160	23.8%	-0.27 [-0.38, -0.15]	•
Subtotal (95% CI)			377			2180	33.8%	-0.15 [-0.53, 0.22]	
Heterogeneity: Tau ² = 0.05;	Chi ² = 1.9	90, df = 1	(P = 0	.17); l² = 47	7%				
Test for overall effect: Z = 0.	.80 (P = 0	.42)							
5.2.3 Other									
Fedders 2019 Face region	173.49	66.3	20	161.1	67.5	20	10.0%	0.18 [-0.44, 0.80]	
Zhang 2013 Inguinal line	3.52	1.32	357	3.45	1.39	2160	23.8%	0.05 [-0.06, 0.16]	
Subtotal (95% CI)			377			2180	33.8%	0.05 [-0.06, 0.17]	•
Heterogeneity: Tau ² = 0.00;	Chi ² = 0.1	l6, df = 1	(P = 0)	.68); l ² = 09	%				
Test for overall effect: Z = 0.	.97 (P = 0	.33)							
Total (95% CI)			825			4439	100.0%	-0.20 [-0.46, 0.05]	•
Heterogeneity: Tau ² = 0.07;	Chi ² = 31	.58. df =	6 (P <	0.0001): l ²	= 81%				
Test for overall effect: Z = 1.		'							-4 -2 0 2
Test for subgroup difference	•		= 2 (P =	: 0 09) l ² =	58.3%				Obese Non-obese

Fig. 5. Meta-analysis for the influence of body mass index (mean differences + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.



Heterogeneity: Tau²=0.0059; H²= 4.093, df= 6.000 (P < 0.001); I²= 75.57% Test for overall effect: Z= 0.0115 (P= 0.991)

Fig. 6. Meta-analysis for the influence of body mass index (correlation coefficients). Abbreviations: RE, random effects; PPT, pressure pain threshold.

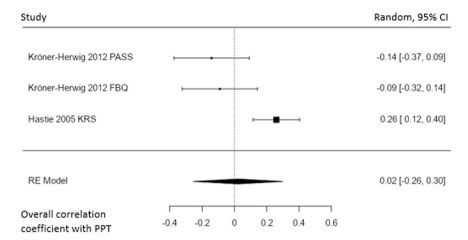
95%CI: -0.32; 0.10). However, two other studies were not included in the meta-analysis, and as a result, qualitative analysis of all five studies reported overall and subgroup-related conflicting results (Table 6). Three studies examined if cognitive impairment influenced PPT. One study showed that having cognitive impairment may result in lower PPT (Girotti et al., 2019), while two other studies (Manning and Fillingim, 2002; Pickering et al., 2020) found no difference (Table 6).

3.11. Comorbidity

Conflicting results. Two studies investigated the effect of comorbidities (Girotti et al., 2019) or previous pain injuries (Manning and Fillingim, 2002) on PPT, and conflicting influence was found (Table 6). Having more comorbidities resulted in a lower PPT (Girotti et al., 2019), but having previous pain injuries resulted in no influence (Manning and Fillingim, 2002).

3.12. Education

Conflicting results. Three studies investigated the effect of education level on PPT, of which one study found that higher education resulted in lower PPT (Zhang et al., 2013), while the other two found no effect (De Rui et al., 2015; Girotti et al., 2019) (Table 6).



Heterogeneity: Tau²=0.0491; H²= 5.702, df= 2.000 (P < 0.003); I²= 82.46% Test for overall effect: Z= 0.156 (P= 0.876)

Fig. 7. Meta-analysis for the Influence of pain vigilance (correlation coefficients). Abbreviations: RE, random effects; PPT, pressure pain threshold.

Study	1	Random, 95% CI
Campbell 2010 CSQ	 i	-0.40 [-0.62, -0.18]
Campbell 2010 PCS	+ +	0.15 [-0.07, 0.37]
Hastie 2005 CSQ	⊢− ∎−−4	-0.16 [-0.30, -0.01]
Kröner-Herwig 2012 PCS	·	-0.02 [-0.25, 0.21]
RE Model		-0.11 [-0.32, 0.10]
Overall correlation coefficient with PPT	-0.8 -0.6 -0.4 -0.2 0 0.2 0.4	

Heterogeneity: Tau²=0.0528; H²= 6.153, df= 3.000 (P < 0.001); I²= 83.75% Test for overall effect: Z= -0.250 (P= 0.803)

Fig. 8. Meta-analysis for the influence of pain catastrophizing (correlation coefficients). Abbreviations: RE, random effects; PPT, pressure pain threshold.

3.13. Ethnicity

Conflicting results. One study reported a difference between two ethnicities (Yang et al., 2013) regarding PPT and two other studies did not (Dawson and List, 2009; Komiyama et al., 2007). However, different ethnicities were included in the three studies (Table 6).

3.14. Gender

Influencing factor. Twenty-eight studies examined the influence of gender on PPT, Fig. 9 shows the meta-analysis, including 19 studies (Garcia et al., 2007; Girotti et al., 2019; Jones et al., 2016; Lautenbacher et al., 2005; Manning and Fillingim, 2002; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Lemming et al., 2015; Matos et al., 2011; Petersen et al., 1992; da Silva et al., 2014; Vatine et al., 1993). Being a woman probably results in having a lower PPT compared to men (SMD: 0.57, 95%CI: 0.39; 0.75). Subgroup analyses of lower arm/hand region (Girotti et al., 2019; Lautenbacher et al., 2005; Moore et al., 2013; Pickering et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 2017; Otto and Dougher, 1985; Chesterton et al., 2002; Shiro et al., 200

2003; Komiyama and De Laat, 2005; Komiyama et al., 2007; Petersen et al., 1992), and face region (Komiyama and De Laat, 2005; Komiyama et al., 2007; Lee et al., 1994; Matos et al., 2011; Petersen et al., 1992; Vatine et al., 1993) also revealed a similar influence of gender (Fig. 9, SMD: 0.68, 95%CI: 0.52; 0.84; and SMD: 0.58, 95%CI: 0.36; 0.79, respectively). Fig. 10 shows the funnel plot with a symmetrical shape, meaning the chance for small study effects is low (Sterne et al., 2011). Of the nine remaining studies, four also reported that being a women resulted in having a lower PPT (De Rui et al., 2015; Kröner-Herwig et al., 2012; Lee et al., 1994; Petrini et al., 2015), but five reported no influence (Dawson and List, 2009; Isselée et al., 2001; Yang et al., 2013, 2014; Bajaj et al., 2001), as such conflicting results regarding qualitative analysis of these nine remaining studies were found (Table 6).

Non-influencing factor: PPT measured at the leg region. Six studies examined the influence of gender on PPT measured at the leg region, Fig. 9 shows the meta-analysis of five studies (Holmgaard et al., 2017; Jones et al., 2016; Manning and Fillingim, 2002; Lemming et al., 2015; Vatine et al., 1993). Gender may have no influence on PPT measured at the leg region (SMD 0.71, 95%CI: -0.22; 1.65). The remaining study also revealed no influence on PPT (Bajaj et al., 2001) (Table 6).

Study or Subgroup	Mean	Men SD	Total	W Mean	omen SD	Total	Weight	Std. Mean Difference IV, Random, 95% CI	Std. Mean Difference IV, Random, 95% Cl
.2.1 Lower arm/hand region	cull	00	. 5141	mourr	00		anaight		
Chesterton 2003 Thenar	42.5	17.8	120	30	14.1	120	3.8%	0.78 [0.51, 1.04]	
Girotti 2019 Carpal joint	40.1	18.9	149	30.6	15.03	206	3.9%	0.57 [0.35, 0.78]	
Komiyama 2005 Thenar	396.1	123	16	261.8	88.9	16	2.3%	1.22 [0.46, 1.98]	
Komiyama 2007 Thenar	363.1	111.3	22	269.5	72	22	2.7%	0.98 [0.35, 1.61]	
_autenbacher 2005 Finger	390		20	302	177.19	20	2.7%	0.41 [-0.22, 1.03]	
Moore 2013 Hand	503.06	160.62	70	410.59	140.79	119	3.7%	0.62 [0.32, 0.92]	
	64.1	53.5	40	32.5	20.6	40	3.7%		
Otto 1985 Finger		335			20.0			0.77 [0.32, 1.23]	
Petersen 1992 Finger	1,160		20	982		20 21	2.7%	0.55 [-0.08, 1.18]	
Pickering 2002 Finger	615.5		21	392	162.5		2.6%	1.39 [0.71, 2.07]	
Shiro 2017 ECRB Subtotal (95% CI)	20	13.6	43 521	16.5	12.5	43 627	3.3% 30.9%	0.27 [-0.16, 0.69] 0.68 [0.52, 0.84]	-
	(D 0.45)	12 0.00				021	30.970	0.00 [0.52, 0.64]	•
Heterogeneity: Tau ² = 0.02; Chi ² = 13.31, df = 9 Test for overall effect: Z = 8.22 (P < 0.00001)	(P = 0.15)	; I² = 32%	D						
I.2.2 Shoulder girdle region									
Fedders 2019 Deltoideus	304.48	143.44	20	226.42	87.06	20	2.7%	0.64 [0.01, 1.28]	⊢
Jones 2016 Biceps + Trapezius	4.4	1.8	18	4.7	2.6	17	2.6%	-0.13 [-0.80, 0.53]	— -
Manning 2002 Shoulder girdle region	7.34	3.78	24	5.69	2.15	24	2.9%	0.53 [-0.05, 1.10]	⊢
Vatine 1993 Sternum	5.2	2.2	14	5.3	1.9	10	2.2%	-0.05 [-0.86, 0.77]	
Subtotal (95% CI)			76			71	10.4%	0.29 [-0.09, 0.67]	•
Heterogeneity: Tau² = 0.04; Chi² = 4.05, df = 3 (Test for overall effect: Z = 1.48 (P = 0.14)	P = 0.26);	I² = 26%							
.2.3 Leg region									
Holmgaard 2017 Tibialis Anterior	1,155.4	445.75	30	1,002.5	356.35	30	3.1%	0.37 [-0.14, 0.88]	+
lones 2016 Rectus femoris + Tibialis anterior	8.6	2.6	18	7.8	3.4	17	2.6%	0.26 [-0.41, 0.93]	-+
emming 2014 Tibialis Anterior	592.5	29.5	48	522	27.5	50	3.0%	2.45 [1.93, 2.98]	
Manning 2002 Quadriceps	11.77	3.65	24	10.4	3.85	24	2.9%	0.36 [-0.21, 0.93]	+
/atine 1993 Malleolus	4.6	2.1	14	4.5	1.9	10	2.2%	0.05 [-0.76, 0.86]	
Subtotal (95% CI)			134			131	13.8%	0.71 [-0.22, 1.65]	
Heterogeneity: Tau² = 1.03; Chi² = 48.28, df = 4 Fest for overall effect: Z = 1.50 (P = 0.13)	(P < 0.000	001); l² =	92%						
I.2.4 Face region	404.0	70 70	00	450.0	50.44	00	0.70/	0.401.0.00.4.051	
Fedders 2019 Face region		73.78	20	152.8	59.44	20	2.7%	0.42 [-0.20, 1.05]	
Holmgaard 2017 Temporalis		178.35	30	337.8	102.4	30	3.0%	0.98 [0.44, 1.52]	
Komiyama 2005 Masseter	208.2		16	146.95	39.85	16	2.3%	1.24 [0.48, 2.01]	
Komiyama 2007 Masseter	192.05	57.7	22	141.35	38.15	22	2.7%	1.02 [0.39, 1.65]	
_ee 1994 Face region	290.19	65.16	104	259.78	53.53	103	3.7%	0.51 [0.23, 0.78]	
Matos 2011 Face region	156.1	64.98	15	121.75	36.95	15	2.4%	0.63 [-0.10, 1.37]	<u> </u>
Petersen 1992 Face region	374	74.5	20	370.5	104	20	2.7%	0.04 [-0.58, 0.66]	-+
Shah 2021 Face region	323.79	150.11	109	258.07	110.41	109	3.8%	0.50 [0.23, 0.77]	-
Vatine 1993 Mastoid process	3.4	1.7	14	3.7	15	10	2.2%	-0.03 [-0.84, 0.78]	— <u></u>
Subtotal (95% CI)			350			345	25.6%	0.58 [0.36, 0.79]	•
Heterogeneity: Tau² = 0.04; Chi² = 12.74, df = 8 Test for overall effect: Z = 5.22 (P < 0.00001)	(P = 0.12)	; I² = 37%	D						
1.2.5 Other									
a Silva 2014 Face region + Tibia	269.67	96.97	61	423.04	164.01	65	3.5%	-1.12 [-1.50, -0.75]	
Garcia 2007 Hypothenar + Tibia + Ulna	4,595.72	528.63	12	4,030.13	737.56	18	2.3%	0.83 [0.07, 1.59]	— -
Garcia 2007 Trigger points	3,616.47	652.17	12	3,161.14	718.12	18	2.4%	0.64 [-0.11, 1.39]	+
ee 1994 Neck region	298.95	66.2	104	254.9	55.55	103	3.7%	0.72 [0.44, 1.00]	-
Shah 2021 Head region	524.96		109		127.02	109	3.8%	0.46 [0.19, 0.73]	
Shah 2021 Neck region Subtotal (95% CI)		158.42	109 407	270.33		109 422	3.7% 19.4%	0.68 [0.41, 0.96] 0.35 [-0.23, 0.93]	-
Heterogeneity: $Tau^2 = 0.46$; $Chi^2 = 73.57$, $df = 5$ Test for overall effect: $Z = 1.19$ ($P = 0.24$)	(P < 0.000	001); l² =						. ,	
Fotal (95% CI)			1488			1596	100.0%	0.57 [0.39, 0.75]	•
Heterogeneity: Tau ² = 0.20; Chi ² = 167.36, df = 3	33 (P ~ 0 (00011-12							· · · · · · · · · · · · · · · · · · ·
neterogeneity. 1au ⁻ – 0.20, Offi ⁻ – 167.36, df = 4	JJ (F ≦ U.U	, I ⁻	- 00%						-4 -2 0 2

Fig. 9. Meta-analysis for the influence of gender (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; ECRB, musculus Extensor carpi radialis brevis; Std., standard.

3.15. Gender role

Conflicting results. One study revealed that gender role influenced PPT (Otto and Dougher, 1985), while the other two studies did not (Manning and Fillingim, 2002; Kröner-Herwig et al., 2012) (Table 6).

3.16. Level of physical activity

Conflicting results. Seven studies investigated if the level of physical activity influenced PPT (Andrzejewski et al., 2010; Jones et al., 2016; Manning and Fillingim, 2002; Shiro et al., 2017; Zhang et al., 2013; Lemming et al., 2015; Kuppens et al., 2019). Fig. 11 displays the meta-analysis of three studies (Manning and Fillingim, 2002; Zhang et al., 2013; Lemming et al., 2015) indicating that the level of physical activity may have no overall influence on PPT (SMD: 0.41, 95%CI:

-0.00; 0.83), but may have an influence on PPT measured at the shoulder girdle/upper arm region (SMD: -0.18, 95%CI: -0.27; -0.09). However, as four studies could not be included in the meta-analysis, also interpretation of the qualitative analysis with all studies was necessary: overall, and subgroup-related conflicting results were found (Table 6).

3.17. Menopause

Non-influencing factor. Two studies (Ozasa et al., 2022; Martínez-Jauand et al., 2013) investigated the influence of being in the menopause, and age of onset of menopause on PPT, but qualitative analysis showed that this factor may not influence PPT (Table 6).

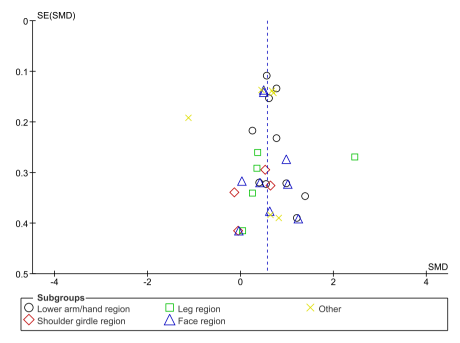


Fig. 10. Funnel plot of gender meta-analysis. Abbreviations: SE, standard error; SMD, standard mean difference.

Higher physical activity				Lower ph	ysical aci	itivity	5	Std. Mean Difference	Std. Mean Difference			
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI			
3.2.1 Leg region												
Lemming 2014 Tibialis anterior	586	19	49	531	20	49	17.0%	2.80 [2.23, 3.36]				
Manning 2002 Quadriceps	10.92	3.5	24	11.25	4	24	17.0%	-0.09 [-0.65, 0.48]				
Zhang 2013 Inguinal line Subtotal (95% CI)	3.34	1.31	726 799	3.49	3.24	1791 1864	24.5% 58.5%	-0.05 [-0.14, 0.03] 0.87 [-0.76, 2.51]				
Heterogeneity: Tau ² = 2.03; Chi ²	= 96.45, df =	2 (P < 0.0	00001); P	² = 98%								
Test for overall effect: Z = 1.05 (F	P = 0.29)											
3.2.2 Arm/shoulder region												
Manning 2002	6.59	3.13	24	6.42	2.8	24	17.0%	0.06 [-0.51, 0.62]				
Zhang 2013 Triceps Subtotal (95% CI)	5.46	1.79	726 750	5.95	2.91	1791 1815	24.5% 41.5%	-0.19 [-0.27, -0.10] -0.18 [-0.27, -0.09]	•			
Heterogeneity: Tau ² = 0.00; Chi ²	= 0.69, df =	1 (P = 0.4	1); l ² = 09	6								
Test for overall effect: Z = 4.14 (F	P < 0.0001)											
Total (95% CI)			1549			3679	100.0%	0.41 [-0.00, 0.83]	◆			
Heterogeneity: Tau ² = 0.18; Chi ²	= 106.81, df	= 4 (P < 0	.00001);	l² = 96%								
Test for overall effect: Z = 1.95 (F	P = 0.05)	•							 -4 -2 0 2 4 Higher physical activity Lower physical acitvity 			
Test for subgroup differences: Cr	ni² = 1.60, df	= 1 (P = 0	.21), l ² =	37.4%					riighei physical activity Lowel physical activity			

Fig. 11. Meta-analysis for the influence of physical activity (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

3.18. Menstrual cycle

Conflicting results. Seven studies investigated the influence of the menstrual cycle on PPT (Alves et al., 2017; Cimino et al., 2000; Isselée et al., 2001; Teepker et al., 2010; Bajaj et al., 2001; Rao et al., 2022; Amodei and Nelson-Gray, 1989), Figs. 12–14 show the meta-analysis of three studies (Teepker et al., 2010; Rao et al., 2022; Amodei and Nelson-Gray, 1989). None of the phases showed differences compared to the other phases of the menstrual cycle regarding PPT (intermenstrual

phase vs. other phases [Fig. 12]: SMD: 1.42, 95%CI: -0.80; 3.63; menstrual phase vs. other phases [Fig. 13]: SMD: -1.01; 95%CI: -2.72, 0.70; premenstrual phase vs. other phases [Fig. 14]: SMD: -0.50, 95% CI: -1.47; 0.47). Regarding the qualitative analysis of all studies, six out of seven studies found an influence of menstrual cycle on PPT, meaning that differences in PPT were found throughout the menstrual cycle. However, the direction differed between studies. As such, the overall and subgroup-related results were conflicting (Table 6).

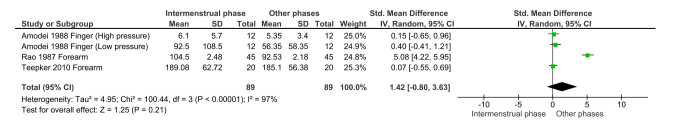


Fig. 12. Meta-analysis for the influence of intermenstrual phase compared to other phases (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

	Menst	trual ph	ase	Othe	r phase	es		Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% Cl
Amodei 1988 Finger (High pressure)	5.9	3.9	12	5.45	4.3	12	24.8%	0.11 [-0.69, 0.91]	
Amodei 1988 Finger (Low pressure)	57.7	51.2	12	73.75	87	12	24.8%	-0.22 [-1.02, 0.59]	
Rao 1987 Forearm	91.06	2.41	45	99.25	2.22	45	25.2%	-3.50 [-4.17, -2.84]	
Teepker 2010 Forearm	169.7	64.11	20	194.79	60.69	20	25.3%	-0.39 [-1.02, 0.23]	
Total (95% CI)			89			89	100.0%	-1.01 [-2.72, 0.70]	
Heterogeneity: Tau ² = 2.90; Chi ² = 68. Test for overall effect: Z = 1.16 (P = 0.1		3 (P < 0.	00001);	l² = 96%	, 0			_	-4 -2 0 2 4 Menstrual phase Other phases

Fig. 13. Meta-analysis for the influence of menstrual phase compared to other phases (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

	Premer	strual ph	nase	Othe	r phase	es		Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Amodei 1988 Finger (High pressure)	4.8	2.9	12	5	4.8	12	23.9%	-0.05 [-0.85, 0.75]	-+-
Amodei 1988 Finger (Low pressure)	55	65.5	12	75.1	79.85	12	23.9%	-0.27 [-1.07, 0.54]	
Rao 1987 Forearm	94	1.95	45	97.78	2.45	45	26.7%	-1.69 [-2.18, -1.21]	
Teepker 2010 Forearm	200.5	58.65	20	194.79	60.69	20	25.6%	0.09 [-0.53, 0.71]	
Total (95% CI)			89			89	100.0%	-0.50 [-1.47, 0.47]	-
Heterogeneity: $Tau^2 = 0.86$; $Chi^2 = 26.0$ Test for overall effect: Z = 1.02 (P = 0.3		P < 0.000	001); l² =	88%				-	-4 -2 0 2 4 Premenstrual phase Other phases

Fig. 14. Meta-analysis for the influence of premenstrual phase compared to other phases (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

	Depres	sed sca	pula	Non-depre	essed sca	ipula	5	Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Azevedo 2008 Upper Trapezius	19	9	26	26.1	9.6	26	70.9%	-0.75 [-1.32, -0.19]	
Lee 2015 Neck/upper back region	3.53	1.15	12	5.37	1.78	12	29.1%	-1.19 [-2.07, -0.31]	
Total (95% CI)			38			38	100.0%	-0.88 [-1.35, -0.40]	•
Heterogeneity: Tau ² = 0.00; Chi ² = 0 Test for overall effect: Z = 3.62 (P =		(P = 0.4	2); l² = 0	1%					-4 -2 0 2 4 Depressed scapula Non-depressed scapula

Fig. 15. Meta-analysis for the influence of scapular position (mean difference + standard deviation). Abbreviations: CI, confidence interval; SD, standard deviation; Random, random effect methods; Std., standard.

3.19. Scapular position

Influencing factor. Two studies evaluated the influence of a depressed scapular position on PPT, with Fig. 15 showing the meta-analysis of those two studies (Azevedo et al., 2008; Lee et al., 2015). Having a depressed scapular position does result in having lower PPT values compared to having a normal scapular position in the neck/shoulder region (SMD: -0.88, 95%CI: -1.35; -0.40).

3.20. Other personal factors - preliminary evidence

Influencing factors. Hand dominance, psychomotor performance, and manual work were each measured by only one study. Therefore, qualitative analysis is preliminary, but may suggest that having a right hand dominance (Petersen et al., 1992) resulted in a higher PPT on the right side, and that having a better psychomotor performance (Pickering et al., 2002), and performing non-manual work (Zhang et al., 2013) resulted in lower PPT (Table 6).

Non-influencing factors. A preliminary qualitative analysis (evidence based on only one study) may show no influence of ADL (Girotti et al., 2019), forward head posture (Kocur et al., 2019), the use of contraceptives (Isselée et al., 2001), hair colour (Holmgaard et al., 2017), functional capacity (Alfieri et al., 2017), sleep (Karmann et al., 2018), polypharmacy (Girotti et al., 2019) and smoking (Zhang et al., 2013) on PPT (Table 6).

3.21. Differences univariate – multivariate analyses

In four (Girotti et al., 2019; Karmann et al., 2018; Yang et al., 2014; Campbell et al., 2015) out of 12 studies, the result of the multivariate analysis was equal to the result of the univariate analysis. However, in eight studies, multivariate analysis differed from univariate analysis: the influence of age (Lautenbacher et al., 2005), alcohol (only one out of two body regions) (Zhang et al., 2013), gender (Pickering et al., 2002; Yang et al., 2013; Komiyama et al., 2007; Kröner-Herwig et al., 2012), ethnicity (Yang et al., 2013; Komiyama et al., 2007), gender role (Otto and Dougher, 1985; Kröner-Herwig et al., 2012) and physical activity (Lemming et al., 2015) did not withstand after correction for different factors (see paragraph '(possible) influencing factors').

3.22. Sensitivity analyses

Sensitivity analyses correcting for studies that did not report full eligibility criteria showed similar results except for the influencing factor age, of which the SMD changed of direction (SMD: 0.06 to -0.45) (Table S2). Sensitivity analyses correcting for meta-analysis model showed no important differences except for the change in SMD direction of the influencing factors age (SMD: 0.06 to -0.29) and level of physical activity (SMD: 0.41 to -0.08), and the amount of SMD change of the influencing factor menstrual cycle (SMD premenstrual phase: -0.50 to -0.75, and SMD intermenstrual phase: 1.21-1.11) (Table S3).

4. Discussion

4.1. Main findings

The goal of this systematic review and meta-analysis was to summarise all studies that had the purpose to explore clinically measurable personal factors which might influence the PPT in healthy people. Only results of univariate analysis (except for the factor 'sleep') were used for

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interpretation.

Influencing factors for PPT overall were blood pressure (very weak conclusion), gender (moderate conclusion) and scapular position (strong conclusion) and subgroup analyses also revealed an influence of age when PPT was measured at the shoulder girdle/upper arm region (weak conclusion), and of gender when PPT was measured at the lower arm/ hand region and face region (moderate conclusion) according to the meta-analytic approach.

Only a *preliminary conclusion of influence* (based on the result of one study) of hand dominance, psychomotor performance and manual work on PPT was found.

Non-influencing factors for PPT were alcohol consumption (weak conclusion), BMI (moderate conclusion) and pain vigilance (very weak conclusion). Subgroup analyses revealed no influence of BMI when PPT was measured at the shoulder girdle/upper arm region, and of gender when PPT was measured at the leg region according to the meta-analytic approach. Depression (moderate conclusion) and menopause (very weak conclusion) and subgroup analysis of BMI when PPT was measured at the face region (weak conclusion) revealed no influence according to the qualitative approach.

Only a *preliminary conclusion of no influence* (based on the result of one study) of affect, fear, ability to dissociate, locus of control ADL, forward head posture, contraceptives, hair colour, function sleep, polypharmacy, and smoking on PPT was found according to the qualitative approach.

Conflicting results were found for all the other personal factors, but further explanation is needed. The overall meta-analytic approach of age, pain catastrophizing, level of physical activity and menstrual cycle showed no influence on PPT, and the subgroup meta-analysis of age measured in the hand region showed influence on PPT (moderate conclusion). However, less than 2/3 of the studies could be implemented in the different meta-analyses and therefore, it seemed also necessary to summarize all studies in a qualitative approach. This resulted in conflicting results (no influence regarding meta-analyses vs. conflicting results regarding qualitative approach).

5. Limitations of the included studies

First, clearly a lot of conflicting results are presented. This can be explained due to the fact that, regarding most influencing factors, metaanalyses were only possible with a subset of the studies. Only a dominant effect according the meta-analysis of blood pressure, gender, scapular position, alcohol consumption and pain vigilance could be presented. For all the other factors, the qualitative approach or a mix of the qualitative and meta-analytic approach is presented. Second, several limitations can be described regarding the included studies. Many studies did not present sufficient data, which makes inclusion in meta-analysis limited. If more studies could be included in the meta-analyses focusing on the influence of age, BMI, pain catastrophizing, level of physical activity and menstrual phase on PPT, maybe less conflicting results will be found. Gender was also the only factor that was measured in more than 10 studies, and as such the only factor for which small study effects could be checked (Sterne et al., 2011) (which was not present). Next, most (42 out of 54) studies performed univariate analyses and did not compensate for confounders, as such, the described (non-)influencing factors are more 'associated' with PPT instead of 'influencing or predicting' PPT (Varga et al., 2020).

Also, important to mention, is the considerable heterogeneity of most meta-analyses. Only the meta-analysis of scapular position had a low heterogeneity (I^2). A part of the overall considerable heterogeneity could be explained by the body region: when accounting for subgroups based on body region, heterogeneity decreased for age with PPT measured at lower arm/hand- or shoulder girdle/upper arm region, BMI with PPT measured at lower arm/hand- or shoulder girdle/upper arm region, gender with PPT measured at face-, lower arm/hand- and shoulder girdle/upper arm region and physical activity with PPT

measured at shoulder girdle/upper arm region with no to moderate heterogeneity. Another explanation could be the measurement method used for measuring the potentially influencing factors regarding blood pressure (systolic, diastolic or mean arterial pressure), BMI (weight and length measured or part of demographic questions), level of physical activity, pain vigilance and pain catastrophizing (different questionnaires). Especially the use of different questionnaires can lead to slight differences, resulting in higher heterogeneity. The difference in cut-off values for age (e.g. older group from 45 (Girotti et al., 2019; da Silva et al., 2014), 50 (Zhang et al., 2013; Lee et al., 1994), 60 (Lautenbacher et al., 2005; Edwards and Fillingim, 2001) or 70 years old (Pickering et al., 2002)) and menstrual phase (e.g. intermenstrual phase between 12th and 16th day (Amodei and Nelson-Gray, 1989) or 15th and 18th day of menstruation (Rao et al., 2022)) for the different groups could also be an explanation. In addition, sensitivity analyses of age (Table S2 and S3), physical activity and menstrual cycle (Table S3) showed non-robustness of their meta-analysis results, indicating these results are sensitive to relatively small changes. Not all studies examining these influencing factors could be implemented in the meta-analysis, and as such, the results of these meta-analyses were not dominant over the results in the qualitative approach. In this way, a correction for the interpretation was already implemented.

5.1. Relation to other reviews and explanations for findings

The overall conflicting result regarding the influence of age is in line with the review of Tumi et al. (Tumi et al., 2017)., as they also reported an inconsistent direction of the influence of age. However, in the meta-analysis of Tumi et al. (Tumi et al., 2017). tentative lower PPT in older compared to younger people were found. This is in contrast with our meta-analysis, which reported no influence of age on PPT. An imported remark is that only four studies were included in their meta-analysis compared to seven studies in ours. The fact that higher PPT was seen in older age when measured in the arm/shoulder region, can be explained by the reduced somatosensory perception due to aging, leading to a loss of nociceptive function and as such reduced sensitivity (Tinnirello et al., 2021). An explanation that this result was only found in the arm/shoulder region could be due to the difference in muscle mass and fat distribution compared to other regions, as previous research indeed showed a difference in PPT at places with extra subcutaneous fat and with little extra fat (Price et al., 2013).

The overall influence of gender is in line with the review of Riley et al (Riley et al., 1998)., showing that men had higher PPT compared to women, and with the review of Racine et al. (Racine et al., 2012)., in which the mediating factors for the influence of gender on PPT were investigated. This difference could be partly explained by hormone differences, as testosterone shows less nociceptive characteristics (Craft, 2007). These hormone differences can also be suggested by findings of six out of seven included studies, in which PPT differences throughout the menstrual cycle were found. However, the direction of influence is inconsistent and therefore the effect is unclear, this is also in line with the conclusions of a previous review (Iacovides et al., 2015). Martin's review (Martin, 2009) found increased pain sensitivity in the intermenstrual phase, but this could not be confirmed for PPT in current review. In addition, Isselée et al (Isselée et al., 2001). found no differences in PPT between women taking contraceptives and women taking no contraceptives. Remarkable, in the leg region, the influence of gender was non-significant. An explanation could be again the difference in muscle mass and fat distribution in the leg, compared to the face or lower arm/hand (Price et al., 2013).

Having a higher blood pressure resulted in higher PPT and could therefore be described as 'blood pressure-related hypoalgesia', which is confirmed by a recent review (Makovac et al., 2020). An explanation for this phenomenon is still unclear, previously it was described that this could be an early sign of a silent asymptomatic myocardial infarct (Ghione, 1996) or that this relation is mediated through endogenous opioids (McCubbin and Bruehl, 1994). However, according to a more recent review, moderating factors for the link between higher blood pressure and higher PPT seems being a woman, when blood pressure is assessed for 24 h ambulatory, when pain stimuli are provided in the arm/leg or mouth/teeth region and when studies did not adjust for confounders. However, future research should examine the underlying factors for this relation (Makovac et al., 2020).

The influence of scapular position on PPT measured at the trapezius muscles can be explained by the fact that the trapezius muscles and brachial plexus are in a more lengthened position when the scapular is depressed. This lengthened position could lead to increased tension, and as such disrupted sarcomeres within the muscles (Kleinrensink et al., 2000). This again can be the cause of higher mechanical hyperalgesia found with PPT measurements (Azevedo et al., 2008; Lee et al., 2015).

The review of Horn-Hofmann et al (Horn-Hofmann et al., 2015). found an overall damping effect of alcohol on pain threshold, tolerance and intensity; and the review of Alabas et al (Alabas et al., 2012). found that gender role was related to pain threshold and tolerance (no separate analysis of which threshold). However, an effect of these two personal factors could not be revealed for PPT with our meta-analysis and qualitative approach, respectively. However, only one study was included regarding gender role in the current review, so caution for the interpretation of the results is needed. An important remark is that the current review and meta-analysis could make conclusions for the influence on PPT separately (instead of combining different pain threshold modalities), which both other reviews (Horn-Hofmann et al., 2015; Alabas et al., 2012) could not.

6. Clinical implications for future research

The influence of many personal factors (ADL, fear, self-efficacy, ability to dissociate, locus of control, contraceptives, hair colour, forward head posture, function, hand dominance, polypharmacy, psychomotor performance, sleep, smoking and manual work) on PPT was only of preliminary strength of recommendation. Therefore, more research regarding whether there is an influence present or not, in combination with an explanation for that (possible) influence is necessary. Future studies should at least present complete absolute data (mean and standard deviation for each group/factor, or correlation coefficient and sample size) or ideally focus on multiple linear regression analysis as statistical analysis, in which the influencing factors can be determined (Schneider et al., 2010). To date, at least blood pressure, gender (PPT measured in lower arm/hand or face region), scapular position (PPT measured in neck/shoulder region), and age (PPT measured in arm/shoulder region) can be considered in research when determining normative values for PPT. This can be used in clinical practice when interpreting sensitivity to pressure using PPT.

7. Strengths and limitations of the review

The strengths of this review include the double-blinded screening in both phases, the RoB assessment, the data extraction, and the fact that this is the first review in the field of influencing factors for PPT that was not restricted to solely one influencing factor. Thereupon, the performance of various subgroup- and different types of meta-analyses (considering mean differences and correlation values) led to stronger conclusions than based on qualitative analysis alone. Although metaanalysis required two different software programs to analyse both subgroup and correlation results.

The current review and meta-analysis should also be considered in the light of some limitations. First, many included studies were found by hand search screening. Despite the non-specification of the influencing factors in the search strategy, many studies were missed. A possible explanation could be the absence of the P-term 'controls' in the search strategy. However, adding this term would led to an overload (+10 000) of hits to screen. Secondly, our inclusion criteria were restricted to clinically measurable personal factors. This means that there is a possibility that other factors, such as environmental factors or genetics, can influence PPT. Thirdly, grey literature search was not performed, as such publication bias cannot be fully excluded. Fourthly, the goal of our systematic review was prognostic, and therefore the QUIPS was chosen to score the RoB. However, most studies only performed univariate analyses in a cross-sectional/ case-control design and received as such higher RoB scores, making the QUIPS tool too strict or too difficult for scoring our included studies. This resulted also in a rather low to moderate Kappa value. However, we tried to compensate for this pitfall by not considering the domains "Study attrition" and "Study confounding" for the overall RoB score when a study design was cross-sectional/ casecontrol. Fifthly, the LoE allocation was performed only by the first author, ideally, a double blind allocation was set up. Finally, our review focused on all personal factors separately and did not focus on the multivariate analyses, despite some factors can be linked to each other and as such be confounding factors (e.g., age after correcting for gender) leading to different results (as seen in eight out of 12 studies that performed multivariate analyses). However, this was the most feasible way to present our findings in order to create a clear overview for the reader.

8. Conclusion

In summary, age (for PPT assessments at shoulder girdle/upper arm region), blood pressure, gender, and scapular position are personal factors that could be considered when determining normative PPT values. Alcohol consumption, BMI, pain vigilance, depression and menopause are personal factors that do not need to be considered. For the influence of other factors there was only preliminary or conflicting evidence, and should be examined further. Caution for interpretation of these results is advised, because of the univariate analysis of most included studies and because many studies were not eligible to include in meta-analyses. Most meta-analyses had considerable heterogeneity, and most conclusions were weak. More research focusing on personal factors, performing adequate statistics and presenting full absolute data is necessary.

Other information

The details of the protocol were prospectively registered at PROS-PERO (registration number 275 191).

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Declaration of interest

None

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.neubiorev.2022.104727.

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