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STUDY PROTOCOL



Efficacy of donepezil for the treatment of oxaliplatin-induced peripheral neuropathy: DONEPEZOX, a protocol of a proof of concept, randomised, triple-blinded and multicentre trial

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Abstract

Background: The use of oxaliplatin in digestive tract cancers could induce severe peripheral neuropathy (OIPN) decreasing the quality of life of patients and survivors. There is currently, no univocal treatment for these peripheral neuropathies. Donepezil, a reversible inhibitor of cholinesterase, used to treat Alzheimer's disease and dementia, is reported to have a good safety profile in humans, and preclinical data have provided initial evidence of its effective-ness in diminishing neuropathic symptoms and related comorbidities in OIPN animal models.

Methods: The DONEPEZOX trial will be a proof-of-concept, randomised, triple-blinded, and multicentre study. It will be the first clinical trial evaluating the efficacy and safety of donepezil for the management of OIPN. Adult cancer survivors with OIPN that report sensory neuropathy according to QLQ-CIPN20 sensory score (equivalence of a grade \geq 2), at least 6 months after the end of an oxaliplatin-based chemotherapy will be included. Eighty patients will be randomly assigned to receive either donepezil or placebo over 16 weeks of treatment. The primary endpoint will be the rate of responders (neuropathic grade decreases according to the QLQ-CIPN20 sensory score) in the donepezil arm. The severity of OIPN will be assessed by the QLQ-CIPN20 sensory scale before and after 16 weeks of treatment. The comparison versus the placebo arm will be a secondary objective. The other secondary endpoints will be tolerance to donepezil, the severity and features of OIPN in each arm before and after treatment, related-comorbidities and quality of life. Fleming's one-stage design will be used for sample size estimation. This design yields a type I error rate of 0.0417 and power of 91% for a responder rate of at least 30% in donepezil arm. A total of 80 randomized patients is planned.

Discussion: This study will allow, in the case of positive results, to initiate a phase 3 randomized and placebo-controlled (primary endpoint) clinical study to assess the therapeutic interest of donepezil to treat OIPN.

Trial registration: NCT05254639, clincialtrials.gov, Registered 24 February 2022.

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Keywords: Pain, Chemotherapy-induced peripheral neuropathy, Anticholinesterase, Donepezil, Oxaliplatin, Study protocol

Background

Chemotherapy-induced peripheral neuropathy (CIPN) is a common adverse effect of neurotoxic anticancer drugs (platinum salts, taxanes, vinca alkaloids, bortezomib, thalidomide), which impacts the patients' quality of life [1]. CIPN occurs in approximately 68% of patients treated with multiple agents [2]. The incidence depends on the chemotherapy regimen, the duration of exposure, and assessment methods [1, 3]. CIPN is the consequence of damage to the peripheral nervous system caused by neurotoxic anticancer agents. These CIPN are mainly characterized by symmetrical sensory disorders with a distal predominance and without truncal or radicular systematization. Most of the symptoms of CIPN correspond to a clinical picture of sensory polyneuropathy, causing loss of sensitivity, paresthesia, dysesthesia, burning sensations, altered fine motor skills and neuropathic pain. About 30% of patients with sensory CIPN also suffer from neuropathic pain [4, 5]. Motor symptoms such as cramping, twitching, muscle atrophy and vegetative disorders can also occur [1]. With cumulative doses, symptoms progress from the ends of the hands and feet to the wrists and ankles and finally the arms and legs [6]. In addition, the perioral or face region, and auditory system can also be affected [7-9].

Oxaliplatin is widely used to treat digestive tract cancers despite being extremely neurotoxic. This neurotoxicity is often dose-limiting and hampers the efficacy of chemotherapy [10]. Oxaliplatin-induced peripheral neuropathy (OIPN) may occur either shortly after oxaliplatin infusion (up to 90% of patients) [2] characterized mainly by acute and transient cold hyperalgesia, or as late-onset chronic cumulative peripheral neuropathy (30–50% of patients) [11]. The resulting neuropathy symptoms can be persistent and definitive (>24 months [11] and>48 months [12, 13]), and associated with psychological distress (anxiety and depression) and a decrease of patients' health-related quality of life (HRQoL) [14].

Although a number of compounds have been investigated to prevent or treat CIPN, none of them are sufficiently effective to be used routinely in clinical practice [15, 16]. Only duloxetine was recommended by ASCO and ESMO for neuropathic pain-related to CIPN [15, 16]. As a result of the lack of clear evidence, the current strategies used to reduce CIPN essentially consist in modifying of the chemotherapy doses and infusion schemes, although sometimes it continues to worsen despite stopping oxaliplatin. This aggravation can be linked to risk factors such as diabetes and excessive alcohol consumption. Little or no follow-up or treatment is offered to patients developing chronic neuropathy after their chemotherapy [10, 13, 17]. This leads to medical wandering, which is very harmful for the patient. Lastly, OIPN is one reason why, despite negative results of the IDEA trial [18], oxaliplatin-based adjuvant treatment of high-risk stage II and low-risk stage III colon cancers is now used for only 3 months as standard of care as compared the previously 6 months of adjuvant treatment.

Donepezil, a reversible inhibitor of cholinesterase, is used to treat Alzheimer's disease and dementia [19]. Donepezil is reported to have a good safety profile in humans [20–22]. In addition, preclinical data have provided initial evidence of its effectiveness in diminishing neuropathic symptoms and related comorbidities in OIPN animal models [23, 24]. The preclinical evidence and the lack of clinical trial data led us to design the first proof of concept study to assess the efficacy and tolerance of donepezil in the treatment of OIPN in cancer survivors.

Preclinical & clinical arguments for using donepezil to treat OIPN

Preclinical works have demonstrated that donepezil was able to prevent and treat tactile allodynia in oxaliplatintreated animals [23, 24]. More precisely, OIPN was responsible of an increase of the cholinergic neurotransmission involving acetylcholine muscarinic M2 receptors (m2AChRs) in the insular cortex of oxaliplatin-treated rats. In the insular cortex, choline (acetylcholine metabolite) concentration was significantly increased and correlated to mechanical allodynia. Moreover, m2AChRs were also up-regulated, and insular micro-injection of oxotremorine (m2AChRs agonist) suppressed mechanical allodynia. Finally, the antineuropathic effect of donepezil was blocked by insular micro-injection of methoctramine (m2AChRs antagonist) [24]. Interestingly, data have highlighted the beneficial effect of donepezil on neuropathic symptoms in different neuropathic pain conditions, likely found in patients affected by CIPN, without significant adverse effects. Likewise, several studies demonstrated that donepezil induces analgesic and neuroprotective effects [25-29]. Recently, a preclinical study demonstrated that donepezil induced an antineuropathic effect in diabetic mice with neuropathic pain [30].

Clinical studies have confirmed the potential antineuropathic effect of donepezil. Boyle et al. [31] showed in healthy volunteers that donepezil (associated with gabapentin) reduced the pain thresholds (better than gabapentin alone) caused by the stimulation of the sural nerve without severe adverse effects. Similarly, this result was also observed in two studies with patients suffering from various neuropathic pain [32, 33]. Finally, a case report demonstrated an analgesic effect of donepezil in a patient with painful Alzheimer's disease [34]. Moreover, donepezil improved depression and anxiety scores and HRQoL in cancer patients treated by opioids [35].

All of these clinical and preclinical data have thus highlighted the potential beneficial effect of donepezil on neuropathic symptoms, without any severe adverse effect. We therefore hypothesized that the use of donepezil could reduce the symptoms of OIPN and limit the decrease in HRQoL and reduce psychological distress (anxiety and depression) in digestive cancer survivors.

Mechanism of CIPN

Better knowledge of the neurotoxicity mechanism induced by anticancer agents is now available. Peripheral neuropathy can result from various causes: damage of the microtubule transport of axons; distal axonal degeneration [36]; morphological change of the nuclei of dorsal root ganglia (DRG) neurons; alteration of mitochondrial function in axons; and intra-axonal accumulation of sodium and calcium by disruption of voltage-gated sodium (Nav) and potassium channels [37]. Nerve damage depends on the anticancer drug administered. Oxaliplatin is known to damage the nuclei of DRG neurons; cause channelopathies of the Nav channel (variation of expression and functionality) and cold receptors of the transient receptor potential family (TRPM8 and TRPA1) [38]. An increase in the glutamate level in the spinal dorsal horn has also been observed in oxaliplatin-treated animals [39], suggesting an involvement of glutamatergic neurotransmission in pain hypersensitivity development induced by oxaliplatin. The chronic sensory form is considered to be induced by morphological and functional changes in the DRG, resulting from the local deposition and accumulation of oxaliplatin. Mitochondrial toxicity could represent an important pathophysiological basis for the chronic neurotoxicity of platinum derivatives [40], as well as satellite glial cell activation in DRG [41].

Current treatment available for CIPN

Several therapeutic agents (antioxidants, neuroprotective agents, analgesics, anticonvulsants, antidepressants, dietary supplements, etc.) have been found promising for CIPN treatment [42]. However, the evidence of their effectiveness remains controversial and the treatment of CIPN is still largely symptomatic. Antidepressants, antiepileptics and current analgesics are the only options available to clinicians, mainly for treating neuropathic pain symptom; however, their effectiveness is limited and associated with significant adverse events like drowsiness. Two studies demonstrated the potential interest of duloxetine for managing CIPN [43, 44], but these studies assessed only the neuropathic pain component (which represents about 30% of patients with CIPN [4, 5]), or they were not randomized or controlled. Lastly, a recent study has shown the preventive effect of Renin-Angiotensin System Inhibitors but this must be confirmed in a randomized controlled trial [45]. The recent update of ASCO and ESMO guidelines for the prevention and treatment of CIPN states that no treatment can be recommended for the prevention of CIPN and only duloxetine can be recommended (level: moderate) for the treatment of neuropathic pain related to CIPN [15, 16]. Therefore, to avoid severe CIPN, dose reduction or discontinuation of the neurotoxic anticancer drug is the only option to limit the progression of CIPN [46, 47].

Donepezil, marketed for the treatment of Alzheimer's disease [48], is known to be neuroprotective [49, 50], anxiolytic and antidepressant [51], with a known safety profile based on Alzheimer patients, and may thus be a suitable candidate for treatment of CIPN.

Methods / design

Study design

The present study is a randomised, triple-blinded (patients, clinicians and pharmacists), and multicentre proof-of-concept trial that assesses the efficacy and safety of donepezil in patients with OIPN. The plan is to include 80 patients from 26 cancer centres and hospitals in France. The duration of the study is 20 weeks for the patients, including 4 weeks for treatment initiation, 12 weeks of stable treatment and 4 weeks of follow-up.

Study objectives

The primary objective is to assess the curative efficacy of donepezil on the severity of OIPN in patients having received and completed oxaliplatin-based chemotherapy indicated for colorectal and pancreatic cancers and eliciting QLQ-CIPN20 sensory score \geq 30 / 100 (equivalent to grade \geq 2 peripheral neuropathy). Efficacy will be based on the percentage of responders (decrease of peripheral neuropathic grade according to the QLQ-CIPN20 sensory score) only in the donepezil arm (see the "Study endpoints" chapter for details).

The secondary objectives are safety (type, intensity, frequency of adverse effects and treatment discontinuation rate due to adverse effects) of donepezil, and its efficacy compared to the placebo arm, for:

- Neuropathic symptoms throughout the study (severity and responders),
- Anxiety and depression,
- OIPN grade,
- Neuropathic pain,
- · Health-related quality of life,
- Patient global impression of change (PGIC),
- Consumption of painkillers.

Inclusion and exclusion criteria

Participants will be adult cancer survivors with OIPN diagnosed for more than 3 months and not relieved by the usual treatments.

Inclusion criteria

 Adult Male / Female who received oxaliplatinebased chemotherapy for colorectal or pancreatic cancers,

 \circ QLQ-CIPN20 sensory score \geq 30 / 100 (equivalent to the National Cancer Institute—Common terminology criteria for adverse events (NCI-CTCAE) v4.0 grade of neuropathy \geq 2),

• Diagnosis of OIPN treated or not by stable antineuropathic treatment (opioids, pregabalin, gabapentin, duloxetine and other antidepressants or anticonvulsants) for at least 1 month,

• Chemotherapy completed for at least 6 months,

 \circ Patients affiliated with the French national health insurance,

• Written informed consent,

• French language comprehension.

Exclusion criteria

• Ongoing cancer,

• Pregnancy or breastfeeding (required contraception),

 Patient with chronic progressive disease inducing pain (excluding OIPN),

 Diabetic patient (excluding non-insulin-treated or insulin-treated diabetes of less than 5 years) or presence of a proven diabetic neuropathy

• Other types of neuropathies,

 \circ ALT / AST more than 3 times the normal value,

• Severe cardiovascular disease, bradycardia

(<55 bpm), heart conduction disorders,

 \circ History or active gastroduodenal ulcer,

 \circ Asthma or obstructive pulmonary disease,

 \circ Allergy to done pezil or piperidine derivatives, Drug interactions: CYP3A4 inhibitors (ketoconazole, itraconazole and erythromycin); CYP2D6 inhibitors (quinidine, fluoxetine) and enzymatic inducers (rifampicin, phenytoin, carbamazepine),
Dependence on alcohol and/or drugs,

Psychotic disorders, patient under antipsychotics.

No therapeutic change will be generated by the protocol; patients will be treated with donepezil or placebo in addition to their current treatment of OIPN, if any. However, no therapeutic change for OIPN or pain will be allowed throughout the study.

Patients can discontinue the study treatment for any of the following reasons: intolerance to donepezil and significant adverse events; and can be withdrawn from analysis for any of the following reasons: modification of the antineuropathic / analgesic therapy (if any) during the study, withdrawal of consent and serious protocol deviation.

Investigational Medicinal Product (IMP) Donepezil Mylan[®]

The study dose of donepezil (5 mg tablet, MYLAN) will be used according to the recommendations by summary of product characteristics (SPC) and will be identical to the dose approved for treating Alzheimer disease and dementia. According to the SPC of donepezil, the dose of 10 mg/day (2 tablets of 5 mg as a single dose) will be reached after 4 weeks at the dose of 5 mg/day. However, if the patient presents efficacy (decrease of OIPN grade) or adverse effects that appear to be related to the study treatment, the investigator will have the option of continuing treatment at a dose of 5 mg/day, until the end of the study.

To improve adherence to intervention protocols, drug tablet return will be counted by pharmacists and during monitoring.

Placebo

The donepezil and the placebo tablets will be in identical opaque blister and will have the same shape, the same colour and the same branding. The placebo will be taken for 16 weeks with the same administration modalities as the donepezil group.

Patients will be given a diary to remind them of their treatment dosage and to indicate any deviations in their intake. Adverse events will be collected by patients throughout the study in their diaries and during the various visits. The type, intensity and potential relationship with the study treatment will be informed.

Treatment should be discontinued if:

- AST or ALT levels \geq 3 times normal values,
- Appearance of severe cardiovascular disease (as determined by clinician), bradycardia (<55 bpm), cardiac conduction disturbances (assessed by electrocardiogram),
- Development of peptic ulcer disease,
- Development of asthma or obstructive lung disease,
- Development of an allergy to donepezil or piperidine derivatives.
- Pregnancy

Investigators will be allowed to unblind without sponsor intervention in case of emergency (e.g. serious adverse event, death, pregnancy).

Study endpoints

Primary endpoint—QLQ-CIPN20 questionnaire (sensory scale)

The EORTC was developed the QLQ-CIPN20 questionnaire as an auto-questionnaire specifically to assess CIPN severity. The QLQ-CIPN20 contains 20 questions about the patients' experience regarding functional limitations and symptoms related to their CIPN. This autoquestionnaire consist of 3 subscales: sensory, motor and autonomic, which permit a comprehensive picture of the characteristics of CIPN. This auto-questionnaire has been previously validated in clinical trials [52, 53]. The literature suggests that the assessment of CIPN symptoms is preferable with Patient-Reported Outcomes (PROs) rather than Clinician-Reported Outcomes (CROs). Moreover, the QLQ-CIPN20 questionnaire is considered as one of the most appropriate outcome measures [54–56].

The sensory subscale assesses various symptoms such as paraesthesia, dysesthesia, numbness and pain; and is more relevant to assess CIPN than the 11-point numerical rating scale (NRS) for pain, which is specific only for pain symptoms which affects only 30% of patients [4, 5]. Other studies have demonstrated that the comparison of QLQ-CIPN20 and the NCI-CTCAE sensory neuropathic grade provides convergent validity [57, 58]. The QLQ-CIPN20 also provided more detailed information, permitting to distinguish more severity degrees of neuropathy, and was more responsive to change over time than the NCI-CTCAE [58]. Finally, as our study will be a multicentre trial, using the QLQ-CIPN20 will make it possible to carefully standardize the assessment of OIPN between centres.

For the primary endpoint (percentage of responders to the treatment), only the sensory scale scores will be used. According to the studies of Alberti et al. [57] and Le-Rademacher et al. [58], the sensory scale score can be correlated with the NCI-CTCAE sensory neuropathy grade (QLQ-CIPN20 scores < 30: grade 1, \ge 30 and \le 40: grade 2, and > 40 / 100: grade 3–4).

A responder will correspond to a patient moving from a score \geq 30 and \leq 40 to a score < 30 / 100 or from a score > 40 to a score \leq 40 / 100.

Because of the absence of preliminary clinical results in the literature about donepezil and CIPN, and the recommendations of international experts (expertise of our project in the framework of a call for projects), we will perform a proof-of concept study to generate preliminary results for a future randomised placebo-controlled phase 3 study if the results are positive. Then, for the primary endpoint, the responder rate will not be compared to placebo, but assessed only for the donepezil arm after 16 weeks of treatment. Donepezil will be considered effective if 7 or more responders are observed in the 33 patients of the donepezil arm (see "statistics" paragraph for details). The responder rate will be compared to the placebo arm only for the secondary endpoint.

In the event of a beneficial therapeutic effect at the end of treatment (significant reduction in neuropathic disorders [reduction of one grade of neuropathy (according to NCI-CTCAE v4.0 or QLQ-CIPN20 score) or -30% of neuropathic pain (according to NRS 0–10 scale), if present, or PGIC score of 6 or 7]), the investigator will be allowed to request a lifting of the treatment blind, to adapt the further management of the patient and/or refer the patient to a specialist (neurologist, pain physician).

Secondary endpoints

The evolution of all the secondary parameters will be studied in an intragroup (before vs. after treatment) and an intergroup (placebo vs. donepezil arm): i) Sensory, motor and autonomic scales from the QLQ-CIPN20 questionnaire (see below); ii) Grading of OIPN by NCI-CTCAE v4.0: this scale will be used to grade the severity of sensory neuropathy from 1 to 4 (1 = Asymptomatic,2 = Moderate symptoms; limiting instrumental of Activities of Daily Living [ADL], 3=Severe symptoms; limiting self-care ADL, 4=Life-threatening consequences; urgent intervention indicated); iii) Assessment of neuropathic pain: monthly assessment of pain felt in the last week by 11-point NRS. This scale is used for adults and children 10 years old or older patient self-reporting of pain (score of 0 = no pain and 10 = worst possible pain). If NRS \geq 4/10, neuropathic pain will be validated by the DN4 interview questionnaire ("Douleur Neuropathique 4") will be mandatory. The interview portion of the DN4 questionnaire is a clinician-administered screening tool for neuropathic pain [59]. The questionnaire includes 7 items (YES or NO, the test is positive for a score $\geq 3/7$). If the patient have a positive DN4 and a pain intensity $\geq 4/10$ (NRS), neuropathic pain will be confirmed and will be characterized by the Neuropathic Pain Symptom Inventory (NPSI) questionnaire. This self-reported questionnaire assesses neuropathic pain symptoms [60], and includes 12 items; iv) HRQoL by the QLQ-C30 questionnaire (EORTC): this auto-questionnaire (EORTC) assesses the HRQoL of cancer patients, and includes five functional scales, three symptom scales, an overall HRQoL scale, and a number of additional elements assessing other symptoms; v) Anxiety and depression by the hospital anxiety and depression scale (HADS) guestionnaire: this auto-questionnaire detects anxiety and depressive disorders, and includes 14 items rated from 0 to 3 (7 questions for anxiety and 7 for depression). The following interpretation is proposed for each of the two total scores: \leq 7: absence of symptomatology; 8 to 10: suggestive symptomatology; ≥ 11 : indicative symptomatology; vi) Patients' Global Impression of Change (PGIC): PGIC [61] assesses the general effectiveness of the treatment according to the impression of patient, with the scores from 1 to 7 (from aggravated to very improve); vii) Safety: the time to treatment failure will be assess. This is the interval between randomization to treatment discontinuation for any reason. The assessment of adverse events (AEs) associated with study treatment will be also assessed by NCI-CTCAE v4.0. Any AEs will be collected daily by the patient and during the monthly visits. AEs will be categorized according to their intensity, treatment-related or not and by type. At last, viii) Assessment of painkiller consumption (if any): drug consumption for OIPN will be assessed throughout the study.

Methodology and study design (Fig. 1)

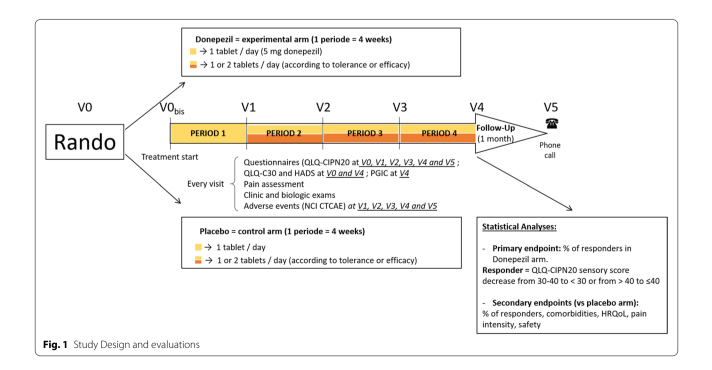
Enrolment

Colorectal and pancreatic cancer survivors having received oxaliplatin-based chemotherapy and eliciting peripheral neuropathy of a grade ≥ 2 (QLQ-CIPN20 sensory scores $\geq 30 / 100$) will be included by clinicians to receive either placebo or donepezil. Patients will either be contacted by phone to propose the study or during one of their consultations at the clinical centre.

Inclusion/randomization visit (V0)

The study will be presented to the patient, along with the objectives, organization, constraints, and assessment questionnaires. If the patient accepts to participate, they will confirm it by signing the informed consent form prior to any study procedure. The following will be performed:

- Validation of patient eligibility (inclusion and noninclusion criteria),
- Clinical examination and body weight,
- Electrocardiogram and heart rate,
- Blood sampling (hepatic and renal function, complete blood count),
- Verification of presence of OPIN (QLQ-CIPN20),
- Randomization,
- Completion of study questionnaires (HADS and QLQ-C30),



- Neuropathic pain assessment: 11-point NRS, and if pain score ≥ 4 / 10 then perform DN4 interview and complete NPSI questionnaire if DN4 interview positive (score ≥ 3 / 7),
- Assessment of painkiller consumption (if applicable),
- Delivery of study treatment according to randomization for a 4-week period: donepezil or placebo.

Visits after 4, 8 and 12 weeks of initiation treatment (V1, V2 and V3)

The following will be performed:

- The patient will return the empty or unused study treatment from the last treatment period,
- Completion of QLQ-CIPN20 questionnaire and 11-point NRS pain,
- Assessment of painkiller consumption (if applicable),
- Electrocardiogram and heart rate,
- Blood sampling (hepatic function, complete blood count),
- Body weight,
- Assessment of safety (adverse effects by NCI CTCAE v4.0),
- Determination of the dose to be delivered according to the treatment efficacy and tolerance, for the 4-weeks period. In the case of efficacy or poor tolerance, the dose will remain at 5 mg/day. If the study treatment is well tolerated but does not show efficacy (according to QLQ-CIPN20 score), the dosage will be increased to 10 mg/day.

End of study treatment visit (V4)

The visit will take place after 16 weeks of treatment or after study treatment discontinuation. The following will be performed:

- The patient will return the empty or unused study treatment from the last treatment period,
- Clinical examination to assess the grade of neuropathy (NCI-CTCAE v4.0),
- Completion of study questionnaires (QLQ-CIPN20, HADS and QLQ-C30),
- Neuropathic pain assessment: 11-point NRS, and if pain score ≥ 4 / 10 then perform the DN4 interview and complete the NPSI questionnaire if the DN4 interview is positive (score ≥ 3 / 7),
- Assessment of painkiller consumption (if applicable),
- Electrocardiogram and heart rate,
- Blood sampling (hepatic function, complete blood count),
- Body weight,

• Assessment of safety (adverse effects by NCI CTCAE v4.0).

Post study treatment follow-up phone call

This phone call will be planned 1-month after the "End of study treatment visit".

The following will be performed during the post-treatment phone call:

- Neuropathic pain assessment: 11-point NRS, and if the pain score ≥ 4 / 10 then the perform DN4 interview and complete the NPSI questionnaire if the DN4 interview positive (score ≥ 3 / 7),
- Completion of study questionnaires (QLQ-CIPN20),
- Assessment of painkiller consumption (if applicable),
- Evaluation of safety (adverse effects by NCI CTCAE v4.0).
- All the questionnaires will be given to the patients during V4 to allow them to complete the questionnaires at home after 1-month. The phone call will remind the patients to return the completed questionnaires to the investigating centre.

Statistical considerations Sample size estimation

Fleming's one-stage design will be used for sample size estimation. Sixty-six evaluable patients (33 by arm) will be accrued. The null hypothesis that the responder rate is 10% (maximal non-efficacy threshold, according to Smith et al. [43]) will be assessed against a one-sided alternative for the donepezil arm. The null hypothesis will be rejected if 7 assessable or more responders are observed in the 33 patients of the donepezil arm. This design yields a type I error rate of 0.0417 and a power of 91% for a responder rate of at least 30% in the donepezil arm (minimal clinical efficacy threshold). Thus, we have decided to include a total of 80 patients (2×40) to take into account lost to follow-up (20%, according to clinical practice with Alzheimer patients, which mainly corresponds to a poor tolerance at 5 mg/day dosage or when switching from 5 mg/day to 10 mg/day dosage), and possible tumor recurrence.

Statistical analysis

All statistical analyses will be performed using SAS software version 9.4 before breaking the randomization codes, according to International Conference on Harmonization-Good Clinical Practice guidelines. A two-sided p-value of less than 0.05 will be considered for the

statistical significance of all the analyses. The results will be expressed as hazard-ratios and 95% confidence intervals. Secondary analyses will be exploratory.

The baseline description of the patients included in the analysis will be carried out, without presenting inferential statistical tests (according to recommendations). As part of a proof-of-concept, only patients who have completed the study (treatment received in its entirety for 16 weeks whatever the dose = per protocol population) will be analyzed for the primary endpoint. The primary endpoint will be the rate of responders and will be described using frequency and percentage. A 95% one-sided confidence interval will also be calculated. Subgroup analyses for the primary endpoint will be planned for each stratum of the randomization (neuropathy grade, age of neuropathy and type of chemotherapy).

For the secondary endpoints, analyses will be performed on the intention-to-treat and per-protocol populations and compared to the placebo arm.

Descriptive statistics will be presented by treatment arms. Continuous variables will be presented as mean and standard deviation (otherwise as median and interquartile range) and will be compared between arms using the unpaired t test or the Mann-Whitney U test when appropriate according to the assumptions of the t-test: (i) the Shapiro-Wilk test will be used to assess normality, and (ii) the Fisher-Snedecor test to assess homoscedasticity. When appropriate (for example, for the sensory scores of the QLQ-CIPN20), these analyses will be supplemented by the recommendations proposed by Vickers and Altman, i.e. ANCOVA, considering baseline scores as covariates. Then, multiple linear mixed models could be run with centers as random-effects. Covariates will be determined according to univariate results and to their clinical relevance, including stratification parameters. The normality of residuals will be studied as described previously. If appropriate, a transformation (for example logarithmic) should be proposed to achieve the normality of dependent variable. The results will be expressed as regression coefficients and 95% confidence intervals. The analyses of other continuous secondary endpoints (11-point pain NRS, DN4 interview, NPSI, HADS and QLQ-C30 questionnaires) will be carried out similarly. For categorical parameters (adverse effects, PGIC, DN4 interview or HADS categorized according to literature), Chi-squared or Fischer exact tests will be used to compare the treatment arms. A generalized linear mixed model (more precisely logistic regression for dichotomous outcomes) will be applied for multivariable analyses. The results will be expressed as relative risks and 95% confidence intervals.

Repeated data collected longitudinally will be analyzed by mixed models (linear or generalized linear according to dependent variable) to study the following fixed effects, arm, time and their interaction taking into account between and within patient variability (patient as random-effect, in addition to center effect) [62]. Particular attention will be paid to safety analysis (by group: type, intensity, frequency of adverse effects). Exit rates due to adverse effects will be considered as a censored data. These criteria will be estimated by the Kaplan-Meier method, compared between groups by the log-rank test, and investigated in a multivariate marginal Cox model considering center effects. The proportionalhazard hypothesis will be verified using Schoenfeld's test and plotting residuals. Finally, to put significant results in perspective, a sensitivity analysis will be conducted to measure the impact of the missing data. A worstcase scenario will be defined as one where patients with missing data do not react to the treatment (whatever the treatment may be). Additional analyses will then be performed according to the statistical nature of missing data (missing at random or not), in particular multiple imputation or estimations proposed by Verbeke and Molen-

Discussion

This study did not encounter any major difficulties in its design and preparation. Nevertheless, there are important points to consider in the design of this type of trial in order to provide a result with an acceptable level of evidence for the scientific and medical community.

berghs [63] adapted specifically to repeated data.

Among these points, we can mention the importance of having a comparison group and at least a double-blind study. Even if in our study, which remains a proof of concept, the comparison with the comparator arm remains a secondary objective, the choice of a placebo comparator group was motivated by three reasons. The first is that there is no active comparator. Indeed, to date, there is no gold standard for the treatment of OIPN [15, 64]. The second reason is the international recommendations of the European Medicines Agency (EMA) for the evaluation of analgesic treatment [65]. These recommendations indicate the need to use a placebo group. The last reason is the placebo effect, which remains very important for pain relief therapies [66]. Thus, the choice to keep a placebo group seemed important to us, and the results of the comparison between the two study arms, even if secondary, will bring a greater level of evidence to our results.

We can also mention the duration of the study. In accordance with the EMA and IMMPACT recommendations, a duration of at least 12 weeks of treatment is recommended to increase assay sensitivity and provide the basis for designing confirmatory trials [67].

It is also important to note the choice of the evaluation criteria. In this study, we chose the sensory score of the QLQ-CIPN20. As previously mentioned, this questionnaire is recommended in several studies for the evaluation of CIPN. Most importantly, this choice also standardizes the assessment of OIPN, thus avoiding measurement bias between centres.

Finally, the study should be as representative as possible of the target population and national clinical practices. For this purpose, 26 investigating centres located throughout France and including academic (regional and university hospitals) and private (cancer centre and private clinic) institutions were selected. It should be noted that a pre-selection of the centres was carried out in order to evaluate their capacity to carry out the study under the conditions of the protocol. The purpose of this selection was to increase the feasibility and success of our study.

Our study also has some limitations. The two main ones being the design in proof of concept and the comparison with the placebo arm only in secondary endpoint. Because of these limitations, the results of this study will have to be confirmed by a randomized, controlled, double-blind study comparing the donepezil arm with the placebo arm as primary endpoint.

Another limitation is the non-inclusion of patients with ongoing cancer. This choice was made because we do not yet have preclinical results on a possible drug interaction between donepezil and the antitumor efficacy of oxaliplatin. Thus, for patient safety, only patients who have completed their chemotherapy will be included. Finally, no electrophysiological or Quantitative Sensory Test exploration is planned in the study. This choice is purely logistical because the vast majority of investigating centres do not have the tools and personnel available to perform this type of exploration.

Abbreviations

AChRs: Acetylcholine Receptors; ADL: Activities of Daily Living; ALT / AST: Alanine aminotransferase / aspartate transaminase; ASCO: American Society of Clinical Oncology; CIPN: Chemotherapy-induced peripheral neuropathy; CROs: Clinician-Reported Outcomes; DN4: "Douleur Neuropathique 4"; DRG: Dorsal root ganglia; EMA: European Medicines Agency; EORTC: European Organisation for Research and Treatment of Cancer; ESMO: European Society for Medical Oncology; HADS: Hospital anxiety and depression scale; HRQoL: Health-related quality of life; IDEA: International duration evaluation of adjuvant therapy; IMMPACT: Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials; Nav: Voltage-gated sodium; NCI-CTCAE: National Cancer Institute—Common terminology criteria for adverse events; NPSI: Neuropathic Pain Symptom Inventory; NRS: Numeric Rating Scale; OIPN: Oxaliplatin-induced peripheral neuropathy; PROs: Patient-Reported Outcomes; PGIC: Patient global impression of change; QLQ-CIPN20: Quality of life questionnaire—Chemotherapy-induced peripheral neuropathy 20; QLQ-C30: Quality of life questionnaire - Cancer 30; SPC: Summary of product characteristics; TRPA1: Transient receptor potential A1; TRPM8: Transient receptor potential M8.

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Data collection and quality management

A clinical research assistant will be dedicated to data entry, coding, security and storage in each centre. Each patient included and the study data will be anonymised. The study data will be collected and managed using an electronic Case Report Form (eCRF, Randomisation Administration Monitoring DAta Management [RAMDAM], Fédération Francophone de Cancérologie Digestive [FFCD]). Randomization will be by minimization according to the following stratification criteria: centre; neuropathy intensity (QLQ-CIPN20 score [\geq 30 to \leq 40] vs [>40 to 100]); and neuropathy age (\leq 1 year vs > 1 year to < 3 years vs > 3 years). A clinical project manager and clinical research assistants will be commissioned by the FFCD (University Hospital of Dijon, Dijon, France) to monitor the progress of the study in accordance with the Standard Operating Procedures implemented by the FFCD and the sponsor (University Hospital of Clermont-Ferrand, Clermont-Ferrand, France), and in accordance with Good Clinical Practices and current French laws. A project manager will be commissioned by the promotor (University Hospital of Clermont-Ferrand) to follow-up and check the global logistics of the study.

Authors' contributions

Nicolas KERCKHOVE wrote the article. NK, DB, DP, DT, KLM, MP, CL and MM led and contributed to the conceptualisation, design and implementation of this research protocol. BP and KLM led the development of the statistical analysis plan. All the authors have read and approved the final manuscript.

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Availability of data and materials

The data set will be the property of the University Hospital of Clermont-Ferrand and FFCD. The principal investigator (DP) and the data-manager will have full access to the final data set. The results will be communicated in a peer-reviewed journal, presented at international congresses and completed online on ClinicalTrials.gov. The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate

The protocol has been approved by the competent authority since the 6 January 2022 (ref: MEDAECNAT-2021–11-0013_2021-005326–23; ANSM, Paris, France) and the ethics committee since the 24 January 2022 (ref: CPP 1444 ME1; CPP Ouest 6, Brest, France). Any substantial modification of the protocol and of the informed consent form will be presented to the independent medical ethics committee. The latter and the competent French authority will be informed of the end of the study.

An independent Data Safety Monitoring Board (DSMB) will be set up to guarantee effective protection of the patients, ensure the ethical conduct of the trial, assess the benefit/risk ratio, and supervise the review of the scientific results during and at the end of the trial. The committee will meet annually and at the request of the sponsor. The committee will be independent of the sponsor and will consist of a CIPN neurologist, a donepezil neurologist, an oncologist and a biostatistician.

The study is currently registered on the clinical trials website under the following number: NCT05254639. The protocol has been in version 4 since the 1st February 2022.

According to the French law on biomedical research, written informed consent must be obtained from patients prior to participation in the study. Patients will voluntarily confirm their willingness to participate in the study, after having been informed (in writing and verbally) by the investigators of all the aspects of the study that are relevant to their decision to participate. They will be informed about requirements concerning data protection, good clinical practices and have to agree to direct access to their individual data. The patients will be informed that they are free to withdraw from the study at any time at their own discretion without necessarily giving reasons.

Finally, the informed consent form was reviewed and validated by a patient association (ARCAD Foundation, committee of the League against Cancer – "Ligue contre le Cancer").

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests. MYLAN will not provide donepezil free of charge and will not participate in any stage of the clinical study or analysis of the results.

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References

- Kerckhove N, Collin A, Condé S, Chaleteix C, Pezet D, Balayssac D. Long-Term Effects, Pathophysiological Mechanisms, and Risk Factors of Chemotherapy-Induced Peripheral Neuropathies: A Comprehensive Literature Review, Front Pharmacol. 2017;8:86.
- 2 Seretny M, Currie GL, Sena ES, Ramnarine S, Grant R, MacLeod MR, et al. Incidence, prevalence, and predictors of chemotherapy-induced peripheral neuropathy: A systematic review and meta-analysis. Pain. 2014;155:2461–70.
- 3 Molassiotis A, Cheng HL, Lopez V, Au JSK, Chan A, Bandla A, et al. Are we mis-estimating chemotherapy-induced peripheral neuropathy? Analysis of assessment methodologies from a prospective, multinational, longitudinal cohort study of patients receiving neurotoxic chemotherapy. BMC Cancer. 2019;19:132.
- 4 de Carvalho BM, Kosturakis AK, Eng C, Wendelschafer-Crabb G, Kennedy WR, Simone DA, et al. A quantitative sensory analysis of peripheral neuropathy in colorectal cancer and its exacerbation by oxaliplatin chemotherapy. Cancer Res. 2014;74:5955–62.
- 5 García de Paredes ML, Del Moral González F, Martínez Del Prado P, Martí Ciriquián JL, Enrech Francés S, Cobo Dols M, et al. First evidence of oncologic neuropathic pain prevalence after screening 8615 cancer patients Results of the On study. Ann Oncol Off J Eur Soc Med Oncol. 2011;22:924–30.
- 6 Armstrong T, Almadrones L, Gilbert MR. Chemotherapy-induced peripheral neuropathy. Oncol Nurs Forum. 2005;32:305–11.
- 7 Hausheer FH, Schilsky RL, Bain S, Berghorn EJ, Lieberman F. Diagnosis, management, and evaluation of chemotherapy-induced peripheral neuropathy. Semin Oncol. 2006;33:15–49.
- 8 Smith EML, Beck SL, Cohen J. The total neuropathy score: a tool for measuring chemotherapy-induced peripheral neuropathy. Oncol Nurs Forum. 2008;35:96–102.
- 9 Giraudet F, Selvy M, Kerckhove N, Pereira B, Barreau F, Nguyen D, et al. Relation between auditory difficulties and bortezomib-induced peripheral neuropathy in multiple myeloma: a single-center cross-sectional study. Eur Arch Oto-Rhino-Laryngol Off J Eur Fed Oto-Rhino-Laryngol Soc EUFOS Affil Ger Soc Oto-Rhino-Laryngol Head Neck Surg. 2022. https:// doi.org/10.1007/s00405-021-07234-1.
- 10 Selvy M, Pereira B, Kerckhove N, Busserolles J, Farsi F, Guastella V, et al. Prevention, diagnosis and management of chemotherapy-induced peripheral neuropathy: a cross-sectional study of French oncologists'

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professional practices. Support Care Cancer. 2021. https://doi.org/10. 1007/s00520-020-05928-6.

- 11 Park SB, Lin CSY, Krishnan AV, Goldstein D, Friedlander ML, Kiernan MC. Long-term neuropathy after oxaliplatin treatment: challenging the dictum of reversibility. Oncologist. 2011;16:708–16.
- 12 André T, Boni C, Navarro M, Tabernero J, Hickish T, Topham C, et al. Improved overall survival with oxaliplatin, fluorouracil, and leucovorin as adjuvant treatment in stage II or III colon cancer in the MOSAIC trial. J Clin Oncol Off J Am Soc Clin Oncol. 2009;27:3109–16.
- 13 Selvy M, Pereira B, Kerckhove N, Gonneau C, Feydel G, Pétorin C, et al. Long-Term Prevalence of Sensory Chemotherapy-Induced Peripheral Neuropathy for 5 Years after Adjuvant FOLFOX Chemotherapy to Treat Colorectal Cancer: A Multicenter Cross-Sectional Study. J Clin Med. 2020;9:2400.
- 14 Mols F, Beijers T, Vreugdenhil G, van de Poll-Franse L. Chemotherapyinduced peripheral neuropathy and its association with quality of life: a systematic review. Support Care Cancer. 2014;22:2261–9.
- 15 Loprinzi CL, Lacchetti C, Bleeker J, Cavaletti G, Chauhan C, Hertz DL, et al. Prevention and Management of Chemotherapy-Induced Peripheral Neuropathy in Survivors of Adult Cancers: ASCO Guideline Update. J Clin Oncol. 2020;38(28):3325–48 (JCO2001399).
- 16 Jordan B, Margulies A, Cardoso F, Cavaletti G, Haugnes HS, Jahn P, et al. Systemic anticancer therapy-induced peripheral and central neurotoxicity: ESMO-EONS-EANO Clinical Practice Guidelines for diagnosis, prevention, treatment and follow-up. Ann Oncol Off J Eur Soc Med Oncol. 2020;31:1306–19.
- 17 Selvy M, Kerckhove N, Pereira B, Barreau F, Nguyen D, Busserolles J, et al. Prevalence of Chemotherapy-Induced Peripheral Neuropathy in Multiple Myeloma Patients and its Impact on Quality of Life: A Single Center Cross-Sectional Study. Front Pharmacol. 2021;12:637593.
- 18 André T, Meyerhardt J, Iveson T, Sobrero A, Yoshino T, Souglakos I, et al. Effect of duration of adjuvant chemotherapy for patients with stage III colon cancer (IDEA collaboration): final results from a prospective, pooled analysis of six randomised, phase 3 trials. Lancet Oncol. 2020;21:1620–9.
- 19 Lee J-H, Jeong S-K, Kim BC, Park KW, Dash A. Donepezil across the spectrum of Alzheimer's disease: dose optimization and clinical relevance. Acta Neurol Scand. 2015;131:259–67.
- 20 Hansen RA, Gartlehner G, Webb AP, Morgan LC, Moore CG, Jonas DE. Efficacy and safety of donepezil, galantamine, and rivastigmine for the treatment of Alzheimer's disease: a systematic review and meta-analysis. Clin Interv Aging. 2008;3:211–25.
- 21 Burns A, Gauthier S, Perdomo C. Efficacy and safety of donepezil over 3 years: an open-label, multicentre study in patients with Alzheimer's disease. Int J Geriatr Psychiatry. 2007;22:806–12.
- 22 Jackson S, Ham RJ, Wilkinson D. The safety and tolerability of donepezil in patients with Alzheimer's disease. Br J Clin Pharmacol. 2004;58(Suppl 1):1–8.
- 23 Kawashiri T, Shimizu S, Shigematsu N, Kobayashi D, Shimazoe T. Donepezil ameliorates oxaliplatin-induced peripheral neuropathy via a neuroprotective effect. J Pharmacol Sci. 2019. https://doi.org/10.1016/j.jphs.2019.05.009.
- 24 Ferrier J, Bayet-Robert M, Dalmann R, El Guerrab A, Aissouni Y, Graveron-Demilly D, et al. Cholinergic Neurotransmission in the Posterior Insular Cortex Is Altered in Preclinical Models of Neuropathic Pain: Key Role of Muscarinic M2 Receptors in Donepezil-Induced Antinociception. J Neurosci Off J Soc Neurosci. 2015;35:16418–30.
- 25 Folkesson A, Honoré PH, Andersen LM, Kristensen P, Bjerrum OJ. Low dose of donepezil improves gabapentin analgesia in the rat spared nerve injury model of neuropathic pain: single and multiple dosing studies. J Neural Transm Vienna Austria. 1996;2010(117):1377–85.
- 26 Kimura M, Hayashida K, Eisenach JC, Saito S, Obata H. Relief of hypersensitivity after nerve injury from systemic donepezil involves spinal cholinergic and γ-aminobutyric acid mechanisms. Anesthesiology. 2013;118:173–80.
- 27 Clayton BA, Hayashida K, Childers SR, Xiao R, Eisenach JC. Oral donepezil reduces hypersensitivity after nerve injury by a spinal muscarinic receptor mechanism. Anesthesiology. 2007;106:1019–25.
- 28 Kim SH, Kandiah N, Hsu J-L, Suthisisang C, Udommongkol C, Dash A. Beyond symptomatic effects: potential of donepezil as a neuroprotective agent and disease modifier in Alzheimer's disease. Br J Pharmacol. 2017;174:4224–32.
- 29 Gomaa AA, Makboul RM, El-Mokhtar MA, Abdel-Rahman EA, Ahmed IA, Nicola MA. Evaluation of the neuroprotective effect of donepezil in type 2 diabetic rats. Fundam Clin Pharmacol. 2020. https://doi.org/10.1111/fcp. 12585.

- 30 Atef MM, El-Sayed NM, Ahmed AAM, Mostafa YM. Donepezil improves neuropathy through activation of AMPK signalling pathway in streptozotocin-induced diabetic mice. Biochem Pharmacol. 2019;159:1–10.
- 31 Boyle Y, Fernando D, Kurz H, Miller SR, Zucchetto M, Storey J. The effect of a combination of gabapentin and donepezil in an experimental pain model in healthy volunteers: Results of a randomized controlled trial. Pain. 2014;155:2510–6.
- 32 Kogure T, Sumitani M, Ikegami K, Abe H, Hozumi J, Inoue R, et al. Donepezil, an Acetylcholinesterase Inhibitor, Can Attenuate Gabapentinoid-Induced Somnolence in Patients with Neuropathic Pain: A Retrospective Chart Review. J Pain Palliat Care Pharmacother. 2017;31:4–9.
- 33 Basnet A, Butler S, Honoré PH, Butler M, Gordh TE, Kristensen K, et al. Donepezil provides positive effects to patients treated with gabapentin for neuropathic pain: an exploratory study. Acta Anaesthesiol Scand. 2014;58:61–73.
- 34 Gharaei H, Shadlou H. A brief report on the efficacy of donepezil in pain management in Alzheimer's disease. J Pain Palliat Care Pharmacother. 2014;28:37–9.
- 35 Bruera E, Strasser F, Shen L, Palmer JL, Willey J, Driver LC, et al. The effect of donepezil on sedation and other symptoms in patients receiving opioids for cancer pain: a pilot study. J Pain Symptom Manage. 2003;26:1049–54.
- 36 LaPointe NE, Morfini G, Brady ST, Feinstein SC, Wilson L, Jordan MA. Effects of eribulin, vincristine, paclitaxel and ixabepilone on fast axonal transport and kinesin-1 driven microtubule gliding: implications for chemotherapyinduced peripheral neuropathy. Neurotoxicology. 2013;37:231–9.
- 37 Grisold W, Cavaletti G, Windebank AJ. Peripheral neuropathies from chemotherapeutics and targeted agents: diagnosis, treatment, and prevention. Neuro-Oncol. 2012;14 Suppl 4:iv45-54.
- 38 Argyriou AA, Bruna J, Marmiroli P, Cavaletti G. Chemotherapy-induced peripheral neurotoxicity (CIPN): an update. Crit Rev Oncol Hematol. 2012;82:51–77.
- 39 Ferrier J, Bayet-Robert M, Pereira B, Daulhac L, Eschalier A, Pezet D, et al. A polyamine-deficient diet prevents oxaliplatin-induced acute cold and mechanical hypersensitivity in rats. PLoS ONE. 2013;8:e77828.
- 40 Zheng H, Xiao WH, Bennett GJ. Functional deficits in peripheral nerve mitochondria in rats with paclitaxel- and oxaliplatin-evoked painful peripheral neuropathy. Exp Neurol. 2011;232:154–61.
- 41 Warwick RA, Hanani M. The contribution of satellite glial cells to chemotherapy-induced neuropathic pain. Eur J Pain Lond Engl. 2013;17:571–80.
- 42 Cavaletti G, Marmiroli P. Chemotherapy-induced peripheral neurotoxicity. Nat Rev Neurol. 2010;6:657–66.
- 43 Smith EML, Pang H, Cirrincione C, Fleishman S, Paskett ED, Ahles T, et al. Effect of duloxetine on pain, function, and quality of life among patients with chemotherapy-induced painful peripheral neuropathy: a randomized clinical trial. JAMA. 2013;309:1359–67.
- 44 Yang Y-H, Lin J-K, Chen W-S, Lin T-C, Yang S-H, Jiang J-K, et al. Duloxetine improves oxaliplatin-induced neuropathy in patients with colorectal cancer: an open-label pilot study. Support Care Cancer. 2012;20:1491–7.
- 45 Uchida M, Kawazoe H, Takatori S, Namba H, Uozumi R, Tanaka A, et al. Preventive Effects of Renin-Angiotensin System Inhibitors on Oxaliplatininduced Peripheral Neuropathy: A Retrospective Observational Study. Clin Ther. 2018. https://doi.org/10.1016/j.clinthera.2018.05.011.
- 46 Winer EP, Berry DA, Woolf S, Duggan D, Kornblith A, Harris LN, et al. Failure of higher-dose paclitaxel to improve outcome in patients with metastatic breast cancer: cancer and leukemia group B trial 9342. J Clin Oncol Off J Am Soc Clin Oncol. 2004;22:2061–8.
- 47 Leonard GD, Wright MA, Quinn MG, Fioravanti S, Harold N, Schuler B, et al. Survey of oxaliplatin-associated neurotoxicity using an interview-based questionnaire in patients with metastatic colorectal cancer. BMC Cancer. 2005;5:116.
- 48 Birks J, Harvey RJ. Donepezil for dementia due to Alzheimer's disease. Cochrane Database Syst Rev. 2018;6(6):CD001190.
- 49 Leyhe T, Hoffmann N, Stransky E, Laske C. Increase of SCF plasma concentration during donepezil treatment of patients with early Alzheimer's disease. Int J Neuropsychopharmacol. 2009;12:1319–26.
- 50 Shafie A, Moradi F, Izadpanah E, Mokarizadeh A, Moloudi MR, Nikzaban M, et al. Neuroprotection of donepezil against morphine-induced apoptosis is mediated through Toll-like receptors. Eur J Pharmacol. 2015;764:292–7.
- 51 Papp M, Gruca P, Lason-Tyburkiewicz M, Willner P. Antidepressant, anxiolytic and procognitive effects of rivastigmine and donepezil in the chronic mild stress model in rats. Psychopharmacology. 2016;233:1235–43.

- 52 Postma TJ, Aaronson NK, Heimans JJ, Muller MJ, Hildebrand JG, Delattre JY, et al. The development of an EORTC quality of life questionnaire to assess chemotherapy-induced peripheral neuropathy: the QLQ-CIPN20. Eur J Cancer Oxf Engl. 1990;2005(41):1135–9.
- 53 Lavoie Smith EM, Barton DL, Qin R, Steen PD, Aaronson NK, Loprinzi CL. Assessing patient-reported peripheral neuropathy: the reliability and validity of the European Organization for Research and Treatment of Cancer QLQ-CIPN20 Questionnaire. Qual Life Res Int J Qual Life Asp Treat Care Rehabil. 2013;22:2787–99.
- 54 Beutler AS, Majithia N, Loprinzi CL. The past and future of "reported outcomes" in studies on chemotherapy neuropathy. Ann Oncol Off J Eur Soc Med Oncol. 2017;28:2631–2.
- 55 Park SB, Kwok JB, Asher R, Lee CK, Beale P, Selle F, et al. Clinical and genetic predictors of paclitaxel neurotoxicity based on patient- versus clinicianreported incidence and severity of neurotoxicity in the ICON7 trial. Ann Oncol Off J Eur Soc Med Oncol. 2017;28:2733–40.
- 56 Dorsey SG, Kleckner IR, Barton D, Mustian K, O'Mara A, St Germain D, et al. The National Cancer Institute Clinical Trials Planning Meeting for Prevention and Treatment of Chemotherapy-Induced Peripheral Neuropathy. J Natl Cancer Inst. 2019;111:531–7.
- 57 Alberti P, Rossi E, Cornblath DR, Merkies ISJ, Postma TJ, Frigeni B, et al. Physician-assessed and patient-reported outcome measures in chemotherapy-induced sensory peripheral neurotoxicity: two sides of the same coin. Ann Oncol Off J Eur Soc Med Oncol ESMO. 2014;25:257–64.
- 58 Le-Rademacher J, Kanwar R, Seisler D, Pachman DR, Qin R, Abyzov A, et al. Patient-reported (EORTC QLQ-CIPN20) versus physician-reported (CTCAE) quantification of oxaliplatin- and paclitaxel/carboplatin-induced peripheral neuropathy in NCCTG/Alliance clinical trials. Support Care Cancer Off J Multinatl Assoc Support Care Cancer. 2017;25:3537–44.
- 59 Bouhassira D, Attal N, Alchaar H, Boureau F, Brochet B, Bruxelle J, et al. Comparison of pain syndromes associated with nervous or somatic lesions and development of a new neuropathic pain diagnostic questionnaire (DN4). Pain. 2005;114:29–36.
- 60 Bouhassira D, Attal N, Fermanian J, Alchaar H, Gautron M, Masquelier E, et al. Development and validation of the Neuropathic Pain Symptom Inventory. Pain. 2004;108:248–57.
- 61 Hurst H, Bolton J. Assessing the clinical significance of change scores recorded on subjective outcome measures. J Manipulative Physiol Ther. 2004;27:26–35.
- 62 Bonnetain F, Fiteni F, Efficace F, Anota A. Statistical Challenges in the Analysis of Health-Related Quality of Life in Cancer Clinical Trials. J Clin Oncol Off J Am Soc Clin Oncol. 2016;34:1953–6.
- 63 Geert Verbeke and Geert Molenberghs. Linear Mixed Models for Longitudinal Data. Springer Book. 2000;978-0-387-22775-7.
- 64 Hershman DL, Lacchetti C, Dworkin RH, Lavoie Smith EM, Bleeker J, Cavaletti G, et al. Prevention and management of chemotherapy-induced peripheral neuropathy in survivors of adult cancers: American Society of Clinical Oncology clinical practice guideline. J Clin Oncol Off J Am Soc Clin Oncol. 2014;32:1941–67.
- 65 EMA. Clinical development of medicinal products intended for the treatment pain. European Medicines Agency. 2018. https://www.ema.europa. eu/en/clinical-development-medicinal-products-intended-treatmentpain. Accessed 5 May 2022.
- 66 Turner JA, Deyo RA, Loeser JD, Von Korff M, Fordyce WE. The importance of placebo effects in pain treatment and research. JAMA. 1994;271:1609–14.
- 67 Dworkin RH, Turk DC, Peirce-Sandner S, Burke LB, Farrar JT, Gilron I, et al. Considerations for improving assay sensitivity in chronic pain clinical trials: IMMPACT recommendations. Pain. 2012;153:1148–58.
- 68 Miltenburg NC, Boogerd W. Chemotherapy-induced neuropathy: A comprehensive survey. Cancer Treat Rev. 2014;40:872–82.
- 69 Cavaletti G, Tredici G, Petruccioli MG, Dondè E, Tredici P, Marmiroli P, et al. Effects of different schedules of oxaliplatin treatment on the peripheral nervous system of the rat. Eur J Cancer Oxf Engl. 1990;2001(37):2457–63.
- 70 Renn CL, Carozzi VA, Rhee P, Gallop D, Dorsey SG, Cavaletti G. Multimodal assessment of painful peripheral neuropathy induced by chronic oxaliplatin-based chemotherapy in mice. Mol Pain. 2011;7:29.

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