

University of Groningen



Inhibition of CYP2D6 with low dose (5 mg) paroxetine in patients with high 10hydroxynortriptyline serum levels - a review of routine practice

Jessurun, Naomi; van Puijenbroek, Eugène P; Otten, Leila S; Mikes, Oenone; Vermeulen Windsant, Annemieke; van Marum, Rob J; Grootens, Koen; Derijks, Hieronymus J

Published in: British Journal of Clinical Pharmacology

DOI: 10.1111/bcp.13201

IMPORTANT NOTE: You are advised to consult the publisher's version (publisher's PDF) if you wish to cite from it. Please check the document version below.

Document Version Publisher's PDF, also known as Version of record

Publication date: 2017

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA): Jessurun, N., van Puijenbroek, E. P., Otten, L. S., Mikes, O., Vermeulen Windsant, A., van Marum, R. J., Grootens, K., & Derijks, H. J. (2017). Inhibition of CYP2D6 with low dose (5 mg) paroxetine in patients with high 10-hydroxynortriptyline serum levels - a review of routine practice. *British Journal of Clinical Pharmacology*, *83*(5), 1149-1151. https://doi.org/10.1111/bcp.13201

Copyright

Other than for strictly personal use, it is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license (like Creative Commons).

The publication may also be distributed here under the terms of Article 25fa of the Dutch Copyright Act, indicated by the "Taverne" license. More information can be found on the University of Groningen website: https://www.rug.nl/library/open-access/self-archiving-pure/taverneamendment.

Take-down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Downloaded from the University of Groningen/UMCG research database (Pure): http://www.rug.nl/research/portal. For technical reasons the number of authors shown on this cover page is limited to 10 maximum.



LETTER TO THE EDITOR

Inhibition of CYP2D6 with low dose (5 mg) paroxetine in patients with high 10hydroxynortriptyline serum levels – a review of routine practice

Correspondence Naomi Jessurun, Netherlands Pharmacovigilance Centre Lareb, Goudsbloemvallei 7, 5237 MH 's-Hertogenbosch, The Netherlands. Tel.: +31 (0)73 646 9700; Fax: +31 (0)73 642 6136; E-mail: n.jessurun@lareb.nl

Received 6 July 2016; Revised 28 October 2016; Accepted 13 November 2016

Naomi Jessurun¹, Eugène P. van Puijenbroek^{1,2}, Leila S. Otten³, Oenone Mikes⁴, Annemieke Vermeulen Windsant⁵, Rob J. van Marum^{5,6}, Koen Grootens⁴ and Hieronymus J. Derijks^{5,7}

¹Netherlands Pharmacovigilance Centre Lareb, 's-Hertogenbosch, The Netherlands, ²Department of Pharmacy: Pharmacotherapy and Pharmaceutical Care, University of Groningen, Groningen, The Netherlands, ³Department of Pharmacy, University of Utrecht, Utrecht, The Netherlands, ⁴Reinier van Arkelgroep, 's-Hertogenbosch, The Netherlands, ⁵Jeroen Bosch Hospital, 's-Hertogenbosch, The Netherlands, ⁶Department of General Practice & Elderly Care Medicine/EMGO Institute for Health and Care Research, VU University Medical Center Amsterdam, The Netherlands, and ⁷Utrecht Institute for Pharmaceutical Sciences, Division of Pharmacoepidemiology & Clinical Pharmacology, Utrecht University, Utrecht, The Netherlands

Tables of Links



These Tables list key protein targets and ligands in this article which are hyperlinked to corresponding entries in http://www.guidetopharmacology.org, the common portal for data from the IUPHAR/BPS Guide to PHARMACOLOGY [1], and are permanently archived in the Concise Guide to PHARMACOLOGY 2015/16 [2].

Nortriptyline, a tricyclic antidepressant (TCA) with selective noradrenergic reuptake inhibitor and little anticholinergic characteristics, is metabolized by CYP2D6 to active metabolites, E-10-hydroxy(OH-)nortriptyline and Z-10-hydroxy(OH-)nortriptyline. Severe depression and depression with psychotic features in the elderly are treated with TCAs. Nortriptyline is preferred in the Netherlands, because it causes the least adverse drug reactions [3]. The therapeutic range of nortriptyline serum levels for anti-depressive treatment lies between 50 μ g l⁻¹ and 150 μ g l⁻¹ [4]. The 10-OH-nortriptyline serum level is preferably kept below 200 μ g l⁻¹ as higher levels are associated with increased occurrence of side effects (e.g. increase in QRS duration and Q-Tc intervals) [5]. So far, only one prospective pharmacokinetic study in five healthy volunteering ultra-rapid metabolizers has been published describing the effects of the addition of 10–20 mg paroxetine (for CYP2D6 inhibition) to 50 mg nortriptyline in order to phenoconvert ultra-rapid metabolizers into poor metabolizers [6]. In the Reinier van Arkelgroep (RvA), 's-Hertogenbosch, the Netherlands, a mental health institution, the addition of low dose (5 mg) paroxetine once daily to patients with high 10-OH-nortriptyline (above 200 μ g l⁻¹) serum levels is applied *ad hoc* to carefully lower the 10-OH-nortriptyline level and maintain patients on nortriptyline therapy. The aim of this review of routine practice was to retrospectively assess the pharmacokinetic impact of this once daily 5 mg paroxetine addition on nortriptyline and 10-OH-nortriptyline serum levels in patients with high 10-OH-nortriptyline serum levels.

Patients treated with nortriptyline in the RvA between 1 July 2011 and 1 July 2015 were considered; patients with at least one high 10-OH-nortriptyline serum level and to whom paroxetine 5 mg was precribed for phenoconversion are described. Patients with co-medication that influences CYP2D6 activity (such as bupropion, cinacalcet, fluoxetine, quinidine, duloxetine, sertraline, terbinafine, amiodarone, cimetidine, dexamethasone, rifampin [7]) disregarded. Nortriptyline and unconjugated were 10-OH-nortriptyline were measured in serum bv high-performance liquid chromatography with photodiode array detection. To assess the impact of paroxetine on nortriptyline metabolism, the last nortriptyline and 10-OH-nortriptyline serum levels before, and the first nortriptyline and 10-OH-nortriptyline serum levels after reaching the steady state situation, which is one week after start of paroxetine, were evaluated.

Four patients received 5 mg paroxetine for phenoconversion. Before the start of paroxetine administration, three patients had nortriptyline serum levels in the therapeutic range and one patient had a nortriptyline serum level below the therapeutic range. All patients had 10-OH-nortriptyline serum levels above 200 μ g l⁻¹. After the addition of 5 mg paroxetine, all subsequent nortriptyline serum levels fell within the therapeutic range and three out of four of the subsequent 10-OH-nortriptyline serum levels decreased to below 200 μ g l⁻¹. The effect of the low dose paroxetine on nortriptyline and 10-OH-nortriptyline serum levels are summarized in Table 1 and shown in Figure 1.

This study suggests that the addition of low dose (5 mg) paroxetine to nortriptyline treatment is able to slow down nortriptyline metabolism. The increase in the ratio between nortriptyline/hydroxynortriptyline serum levels after the addition of paroxetine in all patients supports this. The outcomes are comparable with previous research which showed a decrease of 40% in 10-OH-nortriptyline serum levels after addition of paroxetine [6]. However, the retrospective design does have limitations; for example, the relatively small decrease of 10-OH-nortriptyline serum level in patient 1 could not be explained with the retrieved data.

The intentional introduction of a drug-drug interaction to normalize skewed drug metabolism to optimize drug use is well known. Addition of allopurinol to thiopurine use in patients with high thiopurine methyltransferase activity and the addition of ritonavir to lopinavir use are both comparable interventions that are included in standard care [8, 9]. No adverse drug reactions and changes in tolerability are recorded during the addition, and although both paroxetine and nortriptyline inhibit serotonine reuptake, none of the patients reported signs of serotonine syndrome which would be a possible adverse drug interaction. However, the dose of paroxetine is so low that despite the complex metabolism of this drug, with autoinhibition, the 5 mg once daily dosage will not lead to high paroxetine levels or CYP2D6 saturation [6]. According to the outcomes of this study, the addition of 5 mg paroxetine lowers 10-OH-nortriptyline serum levels and may make treatment with nortriptyline possible for patients who have few other treatment options. To further adress these possibilities, the research will be continued in a prospective design.

ur patients
levels in four
line serum
-1)-nortripty
hydroxy(Oł
levels and
yline serum
on nortript
g paroxetine
5 m
Impact of

Table 1

		No. the state of t	Alantainte Alantaintea	onlintaintaine HO	No.4 minter of	OU northinted	0/ docurrent		Before	A 54 or 10 o
Pt F/M	Age Pt F/M (years)	once daily dose (mg)	serum levels before (μg l ⁻¹)	NUTLIFLY OUT IN THE CONTRIPTY IN EVEN SETUR LEVELS SETUR LEVELS $before (\mu g l^{-1})$ before $(\mu g l^{-1})$	Notripuyinte on notripuy serum levels serum levels after $(\mu g l^{-1})$ after $(\mu g l^{-1})$	before $(\mu g I^{-1})$ after $(\mu g I^{-1})$ after $(\mu g I^{-1})$ serum levels serum levels before $(\mu g I^{-1})$	70 uecrease III 70 increase III paroxecure OH- nortriptyline nortriptyline Nortriptyline/ serum levels Serum levels OH-nortriptyline	70 III.crease III paroxecure nortriptyline Nortriptylin serum levels OH-nortript	paroxecure Nortriptyline/ Nortriptyline/ OH-nortriptyline OH-nortriptyline	Alter paroxetine Nortriptyline/ OH-nortriptyline
1	74	75	74	351	117	308	12%	58%	0.2	0.37
2 F	83	50	65	224	98	109	51%	51%	0.29	0.89
3	44	100	56	226	75	137	39%	34%	0.24	0.54
4 F	68	50	43	215	66	77	64%	53%	0.2	0.85
Averages (± s.d.)	Averages 67.25 ± 16.7 (± s.d.)		59.5 ± 13.2	254 ± 64.8	89 ± 23.0	157.8 ± 103.1	$41.5\% \pm 22\%$	$49\%\pm10\%$	0.23 ± 0.04	0.66 ± 0.25







Figure 1

Increase of nortriptyline and decrease of hydroxynortriptyline serum levels ($\mu g l^{-1}$) after the addition of 5 mg paroxetine to the four patients. (Patient 1 had a dose reduction from once daily 75 mg to once daily 40 mg after the second follow-up nortriptyline/OH-nortriptyline serum level)

Competing Interests

There are no competing interests to declare.

References

- **1** Southan C, Sharman JL, Benson HE, Faccenda E, Pawson AJ, Alexander SP, *et al.* The IUPHAR/BPS Guide to PHARMACOLOGY in 2016: towards curated quantitative interactions between 1300 protein targets and 6000 ligands. Nucl Acids Res 2016; 44: D1054–68.
- 2 Alexander SPH, Fabbro D, Kelly E, Marrion N, Peters JA, Benson HE, *et al.* The Concise Guide to PHARMACOLOGY 2015/16: Enzymes. Br J Pharmacol 2015; 172: 6024–109.
- **3** Landelijke stuurgroep Multidisciplinaire richtlijnontwikkeling. Addendum Ouderen bij de Multidisciplinaire Richtlijn Depressie. (version date: 16-9-2008). Available at www.ggzrichtlijnen.nl/ richtlijn/doc/download.php?id=62&bijlage=1 (last accessed 7 June 2016).
- **4** Berm E, Kok R, Hak E, Wilffert B. Relation between CYP2D6 genotype, phenotype and therapeutic drug concentrations among

nortriptyline and venlafaxine users in old age psychiatry. Pharmacopsychiatry 2016; 49: 186–90.

- **5** Schneider LS, Cooper TB, Severson JA, Zemplenyi T, Sloane RB. Electrocardiographic changes with nortriptyline and 10hydroxynortriptyline in elderly depressed outpatients. J Clin Psychopharmacol 1988; 8: 402–8.
- **6** Laine K, Tybring G, Hartter S, Andersson K, Svensson JO, Widen J, *et al.* Inhibition of cytochrome P4502D6 activity with paroxetine normalizes the ultrarapid metabolizer phenotype as measured by nortriptyline pharmacokinetics and the debrisoquin test. Clin Pharmacol Ther 2001; 70: 327–35.
- **7** P450 drug Flockhart Interaction Table. (version date: 2016, access date). Available at http://medicine.iupui.edu/clinpharm/ddis/ main-table/ (last accessed 20 December 2016).
- 8 Goel RM, Blaker P, Mentzer A, Fong SC, Marinaki AM, Sanderson JD. Optimizing the use of thiopurines in inflammatory bowel disease. Ther Adv Chronic Dis 2015; 6: 138–46.
- **9** Huang X, Xu Y, Yang Q, Chen J, Zhang T, Li Z, *et al.* Efficacy and biological safety of lopinavir/ritonavir based anti-retroviral therapy in HIV-1-infected patients: a meta-analysis of randomized controlled trials. Sci Rep 2015; 5: 8528.