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Rapid Metabolization of Protectin D1 by β-Oxidation of Its Polar Head Chain

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\textbf{ABSTRACT}

Protectin D1 (NPD1, PD1) has been proposed to play a key role in the resolution of inflammation. Aside from its \(\omega\)-monohydroxylated metabolite, little has been reported on its metabolic fate. Upon NPD1 incubation in HepG2 cells, LC-MS/MS revealed the formation of two main metabolites, identified as 2,3-dinor-NPD1 and 2,3,4,5-tetranor-NPD1 by comparison with standards obtained through demanding total chemical syntheses. These data represent the first evidence of \(\beta\)-oxidation occurring in specialized pro-resolving mediators (SPMs), and show that the biotransformation of NPD1 by human hepatoma cells is extremely rapid and faster than leukotriene (LTE\(_4\)). Alike LTE\(_4\), the main metabolic process occurs from the polar head chain of NPD1. It may limit NPD1 systemic circulation and prevent its urinary excretion, making difficult its detection and quantitation \textit{in vivo}. Interestingly, tetranor-NPD1, but not dinor-NPD1, maintained the bioactivity of the parent NPD1, inhibiting neutrophil chemotaxis \textit{in vitro}, and neutrophil tissue infiltration \textit{in vivo}.
INTRODUCTION

The neuroprotectin D1 (NPD1) is an omega-3 lipid mediator arising from the docosahexaenoic acid (DHA, C22:6 n-3) by the action of lipoxygenase-like and other enzymatic activities either by transcellular metabolism\(^1\) or in a single cell type \(^2\). This endogenous dihydroxylated and non-cyclic metabolite was shown to possess anti-inflammatory activities, reducing neutrophil infiltration and accumulation. It participates to the active process of the resolution of inflammation, including wound healing. \(^1,3,4\) The various biological properties of NPD1, its biosynthesis, involvement in many inflammatory and/or infectious diseases in cultured models, rodents and human, have been summarized in many recent reviews and article.\(^5-9\)

The emerging role of NPD1 as a critical mediator of the resolution phase of inflammation \(^8,10\) is making the assessment of its plasmatic concentrations as a potential index of \textit{in vivo} formation a very important issue. The literature reflects the extreme difficulty in quantifying Specialized Pro-resolving Mediators (SPMs), in human fluids, mainly the protectin class of compounds that includes NPD1. Several teams provided evidence of the endogenous production of NPD1, both in healthy volunteers \(^11,12\) and patients with peripheral artery disease\(^13\), metabolic syndrome\(^14\), chronic kidney disease\(^15\) or tuberculosis\(^16\). However, the reported circulating quantities in blood \(^12-14,16-18\) (serum, plasma) or urines\(^19\) are extremely low (much lower than for the other SPM), with levels that often do not exceed the limit of quantification (LOQ) even in cases of severe inflammation or high dose omega-3 supplementation.\(^11,14,15,17,20-22\) Given the surprising discrepancy between the low blood levels measured, the high biological efficacy and relevance attributed to NPD1, it might be possible that the NPD1 plasmatic concentrations may not be predictive of the actual production \textit{in vivo}.

Apart from the fact that NPD1 (as most of the SPMs) requires demanding and careful handling as it is prone to isomerize to its E,E,E-isomer (poor chemical stability with light, air, acid pH), NPD1 may be quickly removed from its bioactive site by rapid local and systemic (hepatic) biotransformation.
Beside the previously reported ω-oxidation generating a biologically active omega-hydroxylated derivative (22-OH-NPD1)\textsuperscript{1,23} in human leukocyte, little is known about NPD1 metabolic fate.

Scheme 1 outlines the different putative metabolic pathways for NPD1. According to the major known pathways of dihydroxylated eicosanoid metabolites, \textsuperscript{24-26} the previously reported 22-OH-NPD1 may be oxidized to the corresponding dicarboxylic acid 22-COOH-NPD1, prior to further prospective β-oxidation of the tail chain. Omega-1 hydroxylation reaction (observed in some resolvins \textsuperscript{27}) seems more unlikely as it should provide an isobaric isomer of 22-OH-NPD1, which should have been detected along with 22-OH-NPD1 if present, unless it is rapidly oxidized to the corresponding ketones. Oxidation of one of the allylic alcohols may also take place as observed in resolvin 18-oxo-RvE\textsubscript{1}, maresin 14-oxo-MaR1\textsuperscript{29} and lipoxin 15-oxo-LXA\textsubscript{4}\textsuperscript{30} for instance. Alternatively, the polar head chain of NPD1 may be shortened by two carbon atoms, resulting in a C20 carbon derivative \textsuperscript{7}, without loss of the central E,E,Z-triene unit flanked on both sides by the two hydroxyl groups. Subsequently, the β-oxidation process may go on, starting from the resulting dinor-NPD1 \textsuperscript{7}. As a result, NPD1 may be shortened by four carbon atoms, leading to the tetranor-NPD1 \textsuperscript{8}. 
Scheme 1. Proposed metabolic pathway for Protectin D1

NPD1 may likely undergo hepatic biotransformation by β-oxidation (scheme 1) since its C1-C6 carbon fragment is identical (scheme 2) to the C13-C18 fragment of 18-COOH-19,20-dinor-LTE_4 and of 18-COOH-19,20-dinor-LTB_4, which are converted to 16-COOH-14,15-dihydro-17,18,19,20-tetranor-LTE_3 and 16-COOH-14,15-dihydro-17,18,19,20-tetranor-LTB_4 respectively. More recently, and highly relevant with respect to NPD1, Lawson et al. 34 also showed that the neuroprostane 7-F_4t-NeuroP_5, which shares the same polar head C1-C6 fragment as NPD1, could not be detected in human urine because of its rapid biotransformation into the isoprostane 5-F_3t-IsoP_6 by β-oxidation (Scheme 2).
Scheme 2. Biotransformation of lipids showing a similar γ-unsaturated carboxylated chain

To address the gaps of knowledge in protectin metabolism, we herein investigated NPD1 biotransformation in human HepG2 cells, as a model of potential hepatic metabolism in man. To confirm the identification of the major novel NPD1 metabolites we developed their total synthesis. We established a quantitative LC-MS/MS method to profile NPD1 and its main β-oxidized derivatives, studied the time-course formation of these novel metabolites in HepG2 hepatoma cells in the presence of different concentrations of the parent NPD1 compound, compared it to LTE 4 metabolism and analyzed human urines from healthy subjects for their presence. Biological activity of the novel metabolites of NPD1 was also evaluated in vitro and confirmed in vivo.

RESULTS AND DISCUSSION

We investigated NPD1 metabolism process upon incubation of known quantities of synthetic NPD1 in cultured cells.
Identification of NPD1 metabolites in HepG2 cells

The analysis of cell media after 4 hours of incubation with human hepatoma cell lines (HepG2), was carried out by RP-HPLC coupled to negative ions electrospray MS in a linear ion trap LC/MS, collecting full scan mass spectra, alignment and subtraction of extracted, single m/z chromatograms from HepG2 media obtained either in the presence or in the absence of 13 µM NPD1,

Along with some remaining NPD1 (at m/z 359), three main peaks were observed at the m/z 305, m/z 333 and m/z 375 (Figure 1) consistent with the molecular weight of the carboxylate anion of tetranor-NPD1 8, dinor-NPD1 7 (scheme 1) and 22-OH-NPD1, respectively.

Figure 1. Analysis of the HepG2 cell medium by RP-HPLC/MS using a linear ion trap mass spectrometer with electrospray source. HepG2 cells were incubated for 4 hours without (panel A) or with 13 µM NPD1 (panel B); cell media were extracted and analyzed by RP-HPLC coupled to a linear ion trap mass spectrometer equipped with electrospray ion source. Full scans (m/z 100-1000) were...
collected and single ion chromatograms extracted, over-imposed and used to locate peaks associated with NPD1 incubation. Significant peaks present in NPD1 incubates only are marked with an arrow.

Tandem mass spectrometric analysis of the abovementioned metabolites generated spectra identical to those of tetranor-NPD1 (Figure 2A), dinor-NPD1 (Figure 2B), and NPD1 (data not shown) synthetic standards (see chemical synthesis below), confirming that NPD1 can indeed undergo several rounds of β-oxidation in a model of human liver cells. MS/MS analysis of the ion at m/z 375, generated a spectrum with major fragment ions at m/z 295, 261, 217 and 153, in line with the fragmentation previously reported for 22-OH-NPD1 (see supporting information file, figure S1).

**Figure 2. MS/MS analysis of NPD1 metabolites in HepG2 cell medium.** The MS/MS spectra of compounds found in HepG2 cell medium after 4 hours of incubation with 13 µM of NPD1 were collected by fragmentation of molecular ions at m/z 305.4 (panel A) and m/z 333 (panel B). MS/MS spectra of synthetic tetranor-NPD1 and dinor-NPD1 are shown in insets.

These two novel metabolites of NPD1 (compounds 7 and 8) represent the first evidence of β-oxidation occurring in SPMs: while potential β-oxidation following ω-oxidation was clearly a concern
in the development of analogs of lipoxins, leading to the development of aspirin-triggered lipoxin analogs protected at the omega end, no identification of β-oxidized metabolites was reported to date at either the tail chain or the polar head chain. In the present *in vitro* study, only traces of 22-OH-NPD1 were detected in HepG2 cells, and no peak could be assigned to the corresponding omega-carboxylated derivative (22-COOH-NPD1), suggesting that, contrary to eicosanoids (LTB4, LTE4, LXB4 for instance), the metabolization of the tail chain of NPD1 is not the main metabolic pathway of NPD1.

In our hands, selected ion monitoring for targeting the expected m/z for putative ketone formation (scheme1) did not allow detection of any oxo-protectin derivatives.

**Chemical synthesis of NPD1 and its dinor and tetranor metabolites (7 and 8)**

NPD1 was synthesized at a ten milligram scale in order to add it at several concentrations in cultured cells and monitor its presence or disappearance upon several incubation periods. Since LC-MS/MS analysis of incubated cell media suggested (see above) the formation of 1,2-dinor 7 and 1,2,3,4 tetranor 8 derivatives (which, herein, will be called dinor-NPD1 and tetranor-NPD1 for better convenience), we envisioned their total synthesis. These two targets served as chemical tools for comparison and full elucidation of the structure of the newly observed metabolite peaks in human hepatocyte models. They were also used for the calibration curves required for quantitation in the time-course experiments (see below).

Based on our flexible strategy reported in 2014, the three targets NPD1, its dinor 7 and tetranor 8 may be obtained using two Wittig olefination reactions via an advanced pivotal intermediate 9. This ynediene 9 is a diastereomer of an ynediene that was previously prepared via Sonogashira cross-coupling reaction, Colvin and Bestmann-Ohira alkynylation reactions. Thus, in the synthesis of the iododiene 10, the (S)-butane-triol was simply replaced by its (R)-enantiomer, which can be prepared from (R)-malic acid, much cheaper. The terminal alkyne 11 may be synthesized from (S)-butane-triol via a Bestmann-Ohira
alkynylation reaction and suitable protection-deprotection sequence. The retrosynthetic analysis is depicted in scheme 3.

Scheme 3. Retrosynthetic analysis for NPD1 and its β-oxidized metabolites 7 and 8.

The total synthesis of NPD1 and its dinor 7 and tetranor 8 is illustrated in scheme 4.

The alcohol 9 was oxidized to aldehyde 12 and subsequently converted to aldehyde 13 (previously reported36 with a different configuration for the synthesis of AT-NPD1).
Scheme 4. Total synthesis of NPD1 and its metabolites 7 and 8.

Then, using the same pivotal late advanced key aldehyde 13, the total synthesis of NPD1 and its dinor-NPD1 metabolite 7 was achieved in a total of 27 and 22 steps in 2.84% and 3.92% overall yield with a longest linear sequence respectively. Compared to our previously reported synthesis of the key aldehyde
intermediate 13 yields were slightly improved. Replacement of a TBS group by a TBDPS group facilitated some selective deprotection steps before the Sonogashira reaction leading to ynediene 9.

The synthesis of NPD1-tetranor 8 (scheme 4) was performed using the same strategy as for NPD1 and dinor-NPD1 7. However, the shortening of the polar head made it more difficult to obtain, due to the activated allylic methylene in the alpha-position to the carboxylated function.

The Wittig reaction between the pivotal aldehyde 13 with the (2-carboxyethyl)triphenylphosphonium bromide 14 in the presence of NaHMDS followed by treatment with TMSCHN 2 afforded the desired Wittig product 15 in 69% yield. However, contrary to our previous Wittig reactions with this aldehyde 13, the olefination reaction was not stereoselective, leading to a non-separable Z/E mixture (ratio: 70/30 (NaHMDS as base, -78°C to 0°C, 69%) to 82/18 (n-BuLi, THF:DMSO, -5°C, 71%). Colder temperatures affected the yields without significantly improving the Z/E-stereoselectivity (-15°C, Z/E: 85/15, 55% yield).

Surprisingly, silyl deprotection of ynediene 15 using fluoride anions led to triol 16 instead of the desired diol 17, suggesting an unexpected addition of a hydroxyl group on the isolated Z-double bond. Mild acidic conditions (CSA in MeOH, 0°C) provided the diol 17 (74%).

To overcome the lack of stereoselectivity with the carboxylated phosphonium 14, the carboxylic function was created after the Wittig reaction. Thus, using the THP-protected phosphonium salt 18, aldehyde 13 was converted to the olefin 19 with an excellent Z-selectivity.

Removal of the THP-group was not selective, providing the expected homoallylic alcohol 20 in poor yield (36%) along with partial or full TBS-deprotection. The resulting diol and triol mixture were fully reprotected in the presence of TBSCl and imidazole prior to the selective deprotection of the primary silyl ether (63 % yield over two steps). Conversion of the resulting primary alcohol 20 to aldehyde and carboxylic acid was more difficult than expected. The Dess-Martin reaction and the Pinnick oxidation failed, leading to degradation products only. Oxidation with PDC in DMF afforded the desired
carboxylic function in poor yield (20% yield), albeit complete conversion of alcohol 20. BAIB/TEMPO oxidation\textsuperscript{38} fully converted the alcohol 20 to its corresponding carboxylic acid. However tentative purification by flash chromatography over deactivated silica gel led to polar degradation products. Esterification with TMS-diazomethane was then conducted on the crude product to afford the desired ester 17 in 46 % yield over two steps. As for the carboxylic acid, poor stability is observed.

Subsequently, chemo-selective reduction of the triple bond using Spur protocole\textsuperscript{39} provided the expected E,E,Z-conjugated triene 21 in modest yield (42%). Unfortunately, the compound partially decomposed during the reaction. This unexpected reactivity/stability compared to our previous experience with such reaction\textsuperscript{36, 40} is probably due to the highly activated methylene of the polar chain head. Ester hydrolysis with lithium hydroxide yielded the targeted NPD1-tetranor acid 8 in good yield (88%). Thus, the total synthesis of the tetranor-NPD1 metabolite 8 was achieved in a total of 21 steps and 2.5% overall yield with a longest linear sequence.

**Quantitative determination of NPD1, dinor-NPD1 (7), and tetranor-NPD1 (8)**

The availability of these synthetic standards allowed confirmation of the newly observed metabolites 7 and 8 in HepG2 cells, and the development of a sensitive quantitation method based on RP-HPLC separation coupled to selected reaction monitoring in a triple quadrupole mass spectrometer (Figure S2), using \textit{d}_4-LTB\textsubscript{4}, a tetradeuterated arachidonic acid derivative sharing essential structural features (i.e. a E,E,Z-conjugated triene flanked by two hydroxyls groups) with NPD1 and its β-oxidized metabolites, as internal standard (calibration curves, supporting information file, Figure S3).

Time-course incubation of high concentrations of NPD1 (13 μM) showed the presence of significant amounts of β-oxidized metabolites of NPD1 already after 30 minutes of cell incubation with NPD1, with the amount of the parent compound progressively decreasing and reaching near zero after 24h of
incubation (Figure 3A). Dinor-NPD1 7 relative amount increased up to 8h, but also sharply decreased at 24h, while tetranor-NPD1 represented the main metabolite remaining at 24h. The tetranor-NPD1 steadily increased up to 24h, a time point at which it represented almost 9% of NPD1 measured at time zero (Figure 3B).

Figure 3. Time-course of NPD1, dinor-NPD1 and tetranor-NPD1, after incubation of HepG2 cells with 13 µM NPD1. The medium of HepG2 cells was collected after different incubation times and aliquots were quantitatively analyzed by LC-MS/MS. Panel A: nmoles of NPD1 and its metabolites normalized per mg proteins. Panel B: NPD1, dinor-NPD1 and tetranor-NPD1 nmoles amounts expressed as relative percentage of the amount of NPD1 at T0. Left Y-axis: NPD1 and Right Y-axis: dinor-NPD1 (7) and tetranor-NPD1 (8).

As expected based on the incubations with NPD1, the incubation of HepG2 cells with 13 µM of dinor-NPD1 (7), also led to the formation of tetranor-NPD1 8 (Figure 4), confirming a second round of β-oxidation from dinor-NPD1 7. The amount of dinor-NPD1 expressed in nmoles/mg protein rapidly decreased, and at 24h less than 0.1% was present in the incubation medium. Tetranor-NPD1 steadily
increased reaching a maximum at 8h, but the amounts detected actually declined at 24h (Figure 4), suggesting that further metabolic conversion may be taking place.

Figure 4. Time-course formation of tetranor-NPD1 upon incubation of HepG2 cells with dinor-NPD1. The medium of HepG2 cells was collected after different incubation times and aliquots were quantitatively analyzed by LC-MS/MS. Panel A: compounds are expressed as relative percentage, the sum (nmoles) of the 2 compounds as 100. Panel B: nmoles of dinor-NPD1 and tetranor-NPD1 normalized per mg proteins.

NPD1 has been characterized as a potent autocrine and paracrine anti-inflammatory and cytoprotective agent, active at nanomolar and subnanomolar concentrations, possibly through the G protein-coupled receptor GPR37. Like other autacoids NPD1 is synthesized at the site of action typically by transcellular metabolism. High circulating concentrations of intact NPD1, such as the one used in our above-mentioned in vitro experiments (13 μM) are therefore highly unlikely, and liver uptake may therefore result in very low concentrations of NPD1 in hepatic cells.
Taking advantage of the sensitivity of the LC-MS/MS method set up for the quantitation of NPD1 and its β-oxidized metabolites, we evaluated the metabolism by HepG2 cells using initial lower concentrations of NPD1. Thus, the use of 2.6 µM and 0.52 µM of NPD1 (instead of 13 µM), resulted in a much faster metabolism (Figures 5), with NPD1 rapidly disappearing from incubates (Figure 5D). More than 50% of NPD1 was metabolized within the first 30 minutes of incubation with 0.52 µM of NPD1. These results are pointing out the saturation of the enzymes involved in the β-oxidation of NPD1 occurring at micromolar concentrations of the substrate. Indeed, the estimated half-life of NPD1 in cultured HepG2 cells decreased from over 4h at 13 µM, to 1h at 2.6 µM and ≈ 30 min at 0.52 µM.

Figure 5. Time-course of NPD1, dinor-NPD1 and tetranor-NPD1, after incubation of HepG2 cells with 2.6 µM NPD1 (Panels A and C) and 0.52 µM NPD1 (Panels B and D). The medium of HepG2 cells was collected after different incubation times and aliquots were quantitatively analyzed by LC-MS/MS. Panel A and B: nmol of NPD1 and its metabolites normalized per mg proteins. Panel C and D: NPD1, dinor-NPD1 and tetranor-NPD1 nmoles amounts expressed as relative percentage of the amount of NPD1 at T0. Left Y-axis: NPD1; Right Y-axis: dinor-NPD1 and tetranor-NPD1.
Concomitantly, the concentration of **tetranor-NPD1** 8 reached a maximum after 4h at 2.6 µM and 0.52 µM (after 8h at 13 µM **NPD1**). Subsequently, especially at submicromolar concentrations of **NPD1**, the amounts of **tetranor-NPD1** sharply declined at longer observation times (Figures 5), indicating that the biotransformation of **NPD1** may proceeds beyond the formation of this tetranor metabolite 8.

In support to this assertion, minor amounts of other metabolites were also observed in addition to the two main metabolites 7 and 8 reported above. Based on RP-HPLC retention time and molecular ions (at m/z 279 and m/z 349, Figure 6), they were tentatively identified as **hexanor-NPD1** and **omega-hydroxylated dinor-NPD1**. Although their stereochemistry and configurations cannot be ascertained and although detected in minute quantities only, their presence suggests both a third round of β-oxidation of the polar head of the native **NPD1** together with simultaneous ω-hydroxylation of the tail chain of the newly formed metabolites.
Figure 6. Analysis of the HepG2 cell medium by RP-HPLC/MS using a linear ion trap mass spectrometer with electrospray ionization. HepG2 cells were incubated for 4 hours without (panel A and B, upper tracing) or with 13 μM NPD1 (panel A and B, lower tracing); cell media were extracted and analyzed by RP-HPLC coupled to a linear ion trap mass spectrometer equipped with electrospray ion source. Full scans (m/z 100-1000) were collected and single ion chromatograms extracted, over-imposed and used to locate peaks associated with NPD1 incubation. Significant peaks present in NPD1 incubates only are marked with an arrow.

NPD1 is a fatty acid derivative. Its liver β-oxidation could take place in peroxisomes and/or mitochondria, due to the metabolic interplay between these organelles; although the β-oxidation system in these compartments is chemically very similar, the reactions are catalyzed by different enzymes (encoded by different genes), often being the rate-limiting step of the process. Also, cofactors (NAD⁺, FAD⁺, CoASH, etc), membrane transporters/channels (ABC transporters, CPTI proteins, etc.), and auxiliary enzymes are involved in the complex regulation of β-oxidation and lipid metabolism in peroxisomes and mitochondria, and may affect the rate of this metabolic reaction. The rapid formation of dinor-NPD1 provides another example of the involvement of the 2,4-dienoyl-CoA reductase in the processing of the CoA ester of a polyunsaturated fatty acid-derivative, as previously reported for the formation of 16-COOH-14,15-dihydro-17,18,19,20-tetranor-LTE₄, 16-COOH-14,15-dihydro-17,18,19,20-tetranor-LTB₄, and isoprostane 5-F₃-IsoP; the suggested mechanism depicted in scheme 1 is identical to that observed in the metabolism of polyunsaturated fatty acids containing the nearest double bond an even number of carbon atoms from the acyl-CoA ester.
Comparison of NPD1 and LTE4 metabolism by HepG2 cells

Since no example of β-oxidation has been reported to date in the SPM class of lipid compounds, we compared the efficiency and rate of biotransformation of NPD1 by HepG2 cells with another lipoxygenase-derived polyunsaturated lipid mediator known to undergo β-oxidation *in vivo*.

Thus, we compared the disappearance rate of NPD1 with that of leukotriene E4 (LTE4) which metabolism was found to exclusively occur from the ω-end of the tail chain only. (A direct β-oxidation process from the existing carboxylic acid at C1 in leukotrienes and lipoxines is blocked by the hydroxyl group at position C5).

Interestingly, under the same conditions of incubations as used for NPD1, Figures 7A and 7B showed that LTE4 remained basically unmetabolized up to 4 h after the beginning of the incubation at both concentrations tested, highlighting a definitely faster biotransformation of NPD1 by human hepatoma cells. It is important to note that, prior to the β-oxidation process which shorten the tail chain, LTE4 must firstly be ω-oxidized to the corresponding 20-COOH-LTE4 via ω-hydroxylation, a two-step passage that may represent the rate-limiting step of its conversion. Thus, NPD1 metabolism may also be faster than LTE4 *in vivo*.
Figure 7. Time-course of NPD1 and LTE₄, after incubation of HepG2 cells with 2.6 µM (Panels A) or 0.52 µM (Panels B). The medium of HepG2 cells was collected after different incubation times and aliquots were quantitatively analyzed by LC-MS/MS. NPD1, and LTE₄ nmoles amounts expressed as percentage of the amount of NPD1 or LTE₄ (respectively) at T0.

Analysis of NPD1 and its metabolites in human urines

In vivo production of eicosanoids has long been assessed through the determination of local or hepatic metabolites excreted in urines. In touch with a potentially slower systemic biotransformation, LTE₄ can be found unmodified in healthy human urines and its urinary concentrations are significantly increased in the presence of liver disease.

Unlike LTE₄, we could not detect NPD1 in urines from healthy volunteers although synthetic NPD1 added to urine samples remained intact up to 24h at room temperature and was quantitatively recovered, in either the presence or absence of the antioxidant (105.8±8.9%, mean ± SD in the absence of BHT). Similarly, Lawson et al. reported that 7-F₄t-NeuroP 5, which shares the same γ-unsaturated polar head C1-C6 fragment as NPD1 (scheme 2), could not be detected in human urine too because of its rapid biotransformation into 5-F₃t-IsoP 6 by β-oxidation.

Likewise, its novel β-oxidized metabolites 7 and 8 could not be positively identified in any of the samples tested. The failure in detecting the corresponding tetranor metabolite of NPD1 in human urines may reflect that additional biotransformations may take place, as suggested by Figure 5, where at sub-micromolar concentrations of NPD1, after reaching a maximal concentration, the amounts of tetranor-NPD1 8 declined in HepG2 and nearly fully disappeared at the longest observation time point.

Besides, we also looked for evidence of ω-oxidation of NPD1, as observed for LTE₄ in HepG2 incubation media, but only minor amounts of 22-OH-NPD1 could be observed,
Autacoids such as eicosanoids and SPMs are characterized by potent activities requiring only minute amounts being produced locally and rapidly inactivated both locally and, possibly, systemically. The lack of urinary excretion of intact NPD1, together with the rapid β-oxidation in HepG2 cells suggests that NPD1, in addition to local biotransformation may undergo a relevant hepatic first-pass effect, accounting for significant variability in plasmatic determinations of NPD1.

**Biological activity of newly uncovered β-oxidized metabolites of NPD1**

As previously reported, NPD1 dose dependently inhibited LTB₄-induced chemotaxis of human neutrophils (Figure 8A), a biological activity that is maintained in the omega-hydroxylated metabolite. Interestingly, the tetranor-NPD1 (8) retained the ability of NPD1 to inhibit LTB₄-induced chemotaxis in human PMNL while dinor-NPD1 (7) was ineffective (Figures 8C and 8B, respectively).
Figure 8. Effect of NPD1 (Panel A), dinor-NPD1 (Panel B) and tetranor NPD1 (Panel C) on LTB$_4$-induced chemotaxis. Human PMNL chemotaxis was assessed in a 48-well modified micro chemotaxis chamber, as described in Methods. Lower chamber contained LTB$_4$ (10 nM) and different concentrations of NPD1 and its β-oxidized metabolites or their solvent (EtOH); after 90 min at 37°C, migrated cells were stained and densitometric analysis performed. Data are normalized to negative controls. Original values obtained in sextuplicate are expressed as mean±SEM of 3-4 independent assays. Statistical analysis was carried out with one way ANOVA, followed by Dunnet multiple comparison test. **p<0.01 vs LTB$_4$ alone.

Tetranor-NPD1 (8) bioactivity was also assessed in an in vivo model of acute pulmonary inflammation in mice. Tetranor-NPD1 at the dose of 10 ng/mouse administered intra-tracheally at the same time as LPS decreased by 20% the number of infiltrating neutrophils at 24h (Figure 9).

These results are pointing to a critical role of the core structure of NPD1 (represented by the conjugated E,E,Z-triene flanked by the two hydroxyl groups) for biological activity while the lack of activity by dinor-NPD1 may involve a critical distance of the carboxyl group.

Figure 9. Neutrophils in BAL from lungs of LPS-challenged mice: effect of tetranor-NPD1. Cells recovered from bronchoalveolar lavage (BAL) fluids of control mice and mice challenged with LPS i.t. in the presence or absence of tetranor-NPD1 (10 ng/mouse) were stained for FACS analysis as described
in Methods. Total granulocytes were gated as CD45 \textsuperscript{+}F4/80 \textsuperscript{-} and neutrophil population as Ly6G \textsuperscript{hi} CD11b \textsuperscript{hi}. True count\textsuperscript{®} tubes were used to determine the absolute number of neutrophils. (*) p< 0.05;(**) p< 0.01 vs LPS. Each value represents the mean±SEM (n=3).

CONCLUSION

Time-course incubation of \textbf{NPD1} in HepG2 cells, a model of human hepatocytes, showed (Figures 3, 4 and 5) the sequential appearance of two novel metabolites of \textbf{NPD1} with a concomitant, rapid decrease in \textbf{NPD1} concentrations. A stereoselective synthesis of the two putative main metabolites of \textbf{NPD1} has been achieved. The strategy (about 22 to 25 steps) features Wittig olefinations, alkyne preparation, hydrometalation, Sonogashira cross coupling reactions while the desired configurations of the two hydroxyl groups, on both sides of the conjugated E,E,Z-triene, are introduced thanks to commercially available enantiopure (S)-1,2,4-butanetriol and (R)-malic acid as chiral pools (scheme 4).

LC-MS/MS data of these synthetic samples matched (Figure 2) data obtained upon incubation of HepG2 cells with \textbf{NPD1} or \textbf{dinor-NPD1} \textsuperscript{7}, providing unequivocal evidence that \textbf{NPD1} can be rapidly metabolized \textit{via} subsequent rounds of \( \beta \)-oxidation to newly reported hepatic metabolites \textbf{dinor-NPD1} \textsuperscript{7} and \textbf{tetranor-NPD1} \textsuperscript{8}. \( \beta \)-oxidation did not require the initial \( \omega \)-oxidation to the corresponding dicarboxylated metabolite, as previously observed for \textbf{LTE}_4 or \textbf{LTB}_4, but proceeded directly from the polar head of \textbf{NPD1}. These novel metabolites of \textbf{NPD1} represent the first evidence of \( \beta \)-oxidation occurring in SPMs.

The formation of the omega oxidized product \textbf{22-OH-NPD1} was also observed in trace amounts in HepG2 cells but the corresponding \textbf{22-COOH-NPD1} and oxo-derivatives could not be detected.

We also observed additional metabolites that may contribute to the final disappearance of \textbf{NPD1} from HepG2 incubations; in particular the presence of a putative \textbf{hexanor-NPD1} suggests that \( \beta \)-oxidation of \textbf{NPD1} may proceed beyond the \textbf{tetranor-metabolite} \textsuperscript{8} positively identified, and lead to smaller
fragments contributing to decreasing concentrations of NPD1 as well as of dinor- and tetranor-
metabolites over time.

These results are of importance since many biologists who do not detect NPD1 in vivo tend to conclude that NPD1 is not formed in vivo. Our results highlight that NPD1 may be metabolized at the time the biological tissues or cells are analysed. Together with the lack of urinary excretion of intact NPD1, these results support the notion of an extremely rapid clearance of NPD1 that may account for variable plasmatic concentrations. The determination of the potential hepatic biotransformation pathway of NPD1 presented here is the first, critical step for the identification of biomarkers that could be used as an index of in vivo production of NPD1.

EXPERIMENTAL SECTION

General Information: All reactions were performed under an atmosphere of argon unless otherwise specified. Reactions run under argon were conducted in oven-dried glassware (120 °C, minimum 12 hours). Inert gas was dried by passing it through solid anhydrous calcium sulphate (Drierite). Anhydrous diethyl ether, tetrahydrofuran (THF) and dichloromethane were obtained from a PureSolv™ PS-400 solvent purification system. Acetonitrile, anhydrous toluene and benzene were purchased from Acros or Aldrich. Triethylamine, diisopropylethylamine, pyridine, and hexamethylphosphoramide (HMPA) were freshly distilled over calcium hydride under an argon atmosphere. All commercially available reagents were used without further purification unless otherwise noted. Thin-layer chromatography (TLC) was performed on aluminum pre-coated silica gel plates from Merck, and developed plates were visualized by UV light (254 nm), p-anisaldehyde, or potassium permanganate. Column chromatography was performed using flash chromatography with the indicated eluent on Davisil 40-63 µm silica gel. In some cases, for
flash column chromatography, deactivated silica (prepared by addition of 46 mL of water to 100 g of silica then stirring on rotavapor for 2 h at rt without applying vacuum) was used.

$^1$H NMR spectra (300 MHz, 500 MHz) and $^{13}$C NMR spectra (75 MHz, 125 MHz) were recorded on Bruker AMX300 or Bruker Avance 500 MHz spectrometers respectively. Chemical shifts are reported relative to chloroform ($\delta$ 7.24 ppm) for $^1$H NMR spectra and chloroform ($\delta$ 77.16 ppm) for $^{13}$C NMR spectra. The $^1$H NMR spectra data are presented as follows: chemical shift, multiplicity (s = singlet, d = doublet, dd = doublet of doublet, dt = doublet of triplet, td = triplet of doublet, t = triplet, q = quartet, quin = quintet, m = multiplet, br. = broad), coupling constant(s) in Hertz (Hz), and integration. All the NMR spectra were assigned with the help of 2D NMR techniques (COSY $^1$H-$^1$H, HMQC and HMBC).

Infrared spectra were obtained using a Perkin-Elmer Spectrum One spectrophotometer. They were reported as wavenumber (cm$^{-1}$) of significant peaks. Mass spectra (ESI or APCI) and high-resolution mass spectra (HRMS) were measured at the University of Montpellier 2 on either a Waters SYNAPT G2-Si High Definition mass spectrometer or a micromass Q-TOF mass spectrometer. Maldi-Tof spectra were run on a Bruker Ultraflex III spectrometer in a positive reflectron mode. Two matrixes were used: DCTB (trans-2-[3-(4-tert-Butylphenyl)-2-methyl-2-propenylidene]malononitrile) and Ditranol (1,8,9-antracenetriol), both enriched with sodium salts (addition of trifluoroacetate). UV-Visible spectra were performed on a Cary 100 spectrophotometer. Purity of all final compounds was 95% or higher (estimated by HPLC analysis).

**HepG2 cell incubations.**

HepG2 cells (a human hepatoma cell line) were cultured in minimal essential medium (MEM) with 10% FBS and maintained at 37°C in humidified atmosphere. At the beginning of the experiments the same number of cells (about 300,000) were plated into 60 mm cell culture dishes and incubated with medium without FBS. NPD1 (13, 2.6 or 0.52 µM, final concentrations), dinor-NPD1 7 (13 µM, final
concentration) or the same volume of ethanol (0.4%), were then added to cells for different times of incubation: 30 min, 1h, 4h, 8h and 24h. After incubation both the medium and the cells were collected and stored at -80°C for the analyses. An aliquot of the medium was spiked with $d_7$-LTB$_4$ (purchased from Cayman, 10 ng/sample), centrifuged and extracted on SPE column as previously published$^{55}$; cell pellets were used for protein determination using the Bradford method.

Direct comparison with LTE$_4$ metabolism by HepG2 was carried out using the same conditions, with 2.6 or 0.52 µM final concentrations of both LTE$_4$ and NPD$_1$. An aliquot of the medium collected at 30 min, 1h and 4 h was spiked with $d_7$-LTB$_4$ and $d_5$-LTE$_4$ (purchased from Cayman, 10 ng/sample), centrifuged and extracted on SPE column as previously$^{55}$ published. Results were expressed as % of the amounts measured at T0.

Analysis of HepG2 cell incubations media. RP-HPLC analysis was carried out as previously published with modifications: the column used for separation was an EVO Kinetex C18 (2.1 x 150 mm x 5 µm, Phenomenex), eluted at a flow rate of 0.5 mL/min with a gradient from 90% solvent A (H$_2$O + 0.05 ml/L acetic acid, pH=5.7) to 100% solvent B (65% acetonitrile + 35% methanol) over 30 minutes. The effluent from the HPLC was directly interfaced into the electrospray source of a Linear Ion Trap mass spectrometer LTQ (Thermo Fisher), operated in negative ion, and full scans (m/z 100 to 1000) were collected. Extracted ion chromatograms from samples with and without NPD$_1$ were matched and subtracted. Chromatographic peaks associated with NPD$_1$ incubations were further analyzed collecting MS/MS spectra (m/z 100 to 400 in consideration of the parent ion analyzed).

Chemical procedures.

(5Z,8R,9E,11E,13Z,15S,17Z)-8,15-dihydroxyicosa-5,9,11,13,17-pentaenoic acid (2,3-dinor-NPD$_1$ 7). To a solution of the above-mentioned methyl ester (8 mg, 0.022 mmol) in MeOH (3 mL) was added 1 N LiOH (1.5 mL, 1.5 mmol) at 0 °C. The mixture was stirred at rt for 2.5 h and diluted with McIlvaine’s phosphate buffer$^{57}$ (pH 5.0). The resulting mixture was extracted with EtOAc (2 x 10 mL). The
combined extracts were washed with brine (1 x 10 mL), dried over MgSO$_4$, filtered through filter paper and concentrated to give the desired **1,2-dinor-NPD1** 7 (7.2 mg, 94%). **R$_f$:** 0.62 (EtOAc). **UV:** (EtOH) $\lambda_{max}$ 261.8, 271.4, 282.6 nm. **$^1$H NMR:** (500 MHz, CD$_3$OD): $\delta$ 6.51 (dd, $J = 12.5, 13.2$ Hz, 1H), 6.32-6.20 (m, 2H), 6.07 (t, $J = 11.0$ Hz, 1H), 5.74 (dd, $J = 6.6, 14.6$ Hz, 1H), 5.50-5.41 (m, 3H), 5.40-5.29 (m, 2H), 4.56 (q, $J = 6.2$ Hz, 1H), 4.12 (q, $J = 6.6$ Hz, 1H), 2.41-2.16 (m, 6H), 2.14-2.00 (m, 4H), 1.65 (quin, $J = 7.4$ Hz, 2H), 1.36-1.25 (m, 2H, OH), 0.96 (t, $J = 7.6$ Hz, 3H) ppm. **$^{13}$C NMR:** (125 MHz, CDCl$_3$): $\delta$ 177.8, 138.0, 135.0, 134.8, 134.7, 132.0, 131.4, 130.6, 128.9, 127.1, 125.3, 73.1, 68.6, 36.4, 36.3, 34.6, 27.8, 26.0, 21.7, 14.9 ppm. **MS:** (ESI) m/z: 333.21 (M-H)$^+$, **HRMS:** (ESI) calculated for C$_{20}$H$_{29}$O$_4$ (M-H)$^+$ 333.2066, found 333.2066. **MS:** (ESI) m/z: 357.20 (M+Na)$^+$. **HRMS:** (ESI) calculated for C$_{20}$H$_{30}$O$_4$Na (M+Na)$^+$ 357.2042, found 357.2045.

**41Z,6R,7E,9E,11Z,13S,15Z)**-methyl 6,13-bis(tert-butyldimethylsilyloxy) octadeca-3,7,9,11,15-pentaenumonic acid 1,2,3,4-tetranor-NPD1 (8). To a solution of ester 21 (10.0 mg, 0.0312 mmol) in MeOH (2 mL) was added 1 N LiOH (1.6 mL, 1.6 mmol) at 0 $^\circ$C. The mixture was stirred at rt for 1.5 h and diluted with McIlvaine’s phosphate buffer$^{57}$ (pH 5.0). The resulting mixture was extracted with EtOAc (2 x 10 mL). The combined extracts were washed with brine (1 x 10 mL), dried over MgSO$_4$, filtered through filter paper and concentrated to give desired **1,2,3,4-tetranor-NPD1** 8 (8.43 mg, 88%). **R$_f$:** 0.42 (EtOAc). **UV:** (EtOH) $\lambda_{max}$ 261.2, 271.2, 281.8 nm. **$^1$H NMR:** (500 MHz, CD$_3$OD): $\delta$ 6.52 (dd, $J = 10.4, 13.8$ Hz, 1H), 6.35-6.19 (m, 2H), 6.07 (t, $J = 11.1$ Hz, 1H), 5.76 (dd, $J = 6.4, 14.7$ Hz, 1H), 5.71-5.55 (m, 2H), 5.51-5.41 (m, 1H), 5.40-5.29 (m, 2H), 4.56 (q, $J = 6.3$ Hz, 1H), 4.20-4.12 (m, 1H), 3.08 (d, $J = 6.5$ Hz, 1.67H), 3.02 (d, $J = 6.5$ Hz, 0.33H), 2.40-2.24 (m, 4H), 2.23-2.15 (m, 1H), 2.11-2.01 (m, 3H) 0.96 (t, $J = 7.0$ Hz, 3H) ppm. **$^{13}$C NMR:** (125 MHz, CDCl$_3$): $\delta$ 177.0, 137.8, 134.9, 134.8, 134.7, 131.4, 130.6, 129.4, 128.9, 125.3, 73.1, 68.5, 41.7, 36.4, 34.0, 21.7, 14.6 ppm. **MS:** (ESI) m/z: 305.17 (M-H)$^+$. **HRMS:** (ESI) calculated for C$_{18}$H$_{25}$O$_4$ (M-H)$^+$ 305.1753, found 305.1748.

**41Z,6R,7E,9E,11Z,13S,15Z)**-methyl 6,13-bis(tert-butyldimethylsilyloxy) octadeca-3,7,9,15-tetraen-11-ynoate (Wittig product 15). To a suspension of (2-carboxyethyl)triphenylphosphonium bromide (0.53
g, 1.27 mmol, prior to use, the phosphonium salt was washed with anhydrous benzene [3 x 8 mL] then
dried under vacuum and over P$_2$O$_5$ for 48 h at 90 °C) in anhydrous THF (13 mL) was added NaHMDS
(1.0 mL, 2.0 mmol, 2M solution in Hexane) at -15 °C. The reaction mixture was stirred for further 30 min
at the same temperature, and then further cooled down to -78 °C. In another flask, aldehyde 13 (0.101 g,
0.21 mmol) diluted in THF (8 mL) was cooled down to -78 °C. Then the ylide was cannulated to the
aldehyde solution under argon. Then temperature was raised to 10 °C in 2 h. The reaction mixture was
quenched with sat. NH$_4$Cl (10 mL) and allowed to warm to rt. After extraction with ethyl ether (3 x 30
mL), the combined organic layers were washed with brine (1 x 15 mL), then dried over NaSO$_4$, filtered,
and evaporated under reduced pressure. The resulting carboxylic acid crude was used for next step without
further purification.

Subsequently, (trimethylsilyl)diazomethane (0.2 mL, 0.396 mmol) was added at 0 °C to a solution of the
crude carboxylic acid (0.115 g, 0.21 mmol) in ethyl ether/methanol (9/1, 3 mL). After stirring at rt for 45
min, the volatiles were removed under reduced pressure. Purification by flash chromatography (silica,
cyclohexane/ethyl ether: 98/02) afforded the expected methyl ester 15 (0.078 g, 69%) as a colorless oil.

The Wittig olefination reaction was not selective, providing a 3Z/3E mixture (ca.70/30 to 85/15, estimated
by 1H NMR) $R_f$: 0.81 (EtOAc/cyclohexane: 2/8). **IR (cm$^{-1}$):** 2960, 2933, 2858, 1693, 1473, 1364, 1150,
1162, 973. **$^1$H NMR:** (500 MHz, CDCl$_3$): δ [for the (3Z)-isomer] 6.47 (dd, $J = 10.9$, 15.6 Hz, 1H), 6.16
(dd, $J = 10.9$, 15.2 Hz, 1H), 5.72 (dd, $J = 6.1$, 15.2 Hz, 1H), 5.67-5.60 (m, 1H), 5.59-5.45 (m, 3H), 5.42-
5.34 (m, 1H), 4.44 (td, $J = 1.7$, 6.6 Hz, 1H), 4.18 (q, $J = 6.6$ Hz, 1H), 3.65 (s, 3H), 3.05 (d, $J = 7.0$ Hz,
2H), 2.41 (t, $J = 7.1$ Hz, 2H), 2.30-2.16 (m, 2H), 2.04 (quin, $J = 7.5$ Hz, 2H), 0.94 (t, $J = 7.5$ Hz, 3H), 0.88
(s, 9H), 0.86 (s, 9H), 0.10 (s, 3H), 0.08 (s, 3H), 0.01 (s, 3H), 0.00 (s, 3H) ppm. **$^{13}$C NMR:** (125 MHz,
CDCl$_3$): δ [for the (3Z)-isomer] 172.4, 141.0, 138.8, 134.4, 128.87, 126.3, 124.0, (2C), 124.0, 123.1,
110.9, 93.5, 83.5, 72.4, 63.7, 52.0, 36.8, 36.5, 33.1, 26.0 (2C), 20.9, 18.4, 18.3, 14.4, -4.4 (2C), -4.7, -4.8
ppm. **MS:** (ESI) m/z: 283.17 (M-2TBSOH+H), 415.27 (M-TBSOH+H$^+$), 569.35 (M+Na$^+$). **HRMS:**
(6R,7E,9E,13S,15Z)-methyl 3,6,13-trihydroxyoctadeca-7,9,15-trien-11-ynoate (Unexpected triol 16). Tetra-n-butylammonium fluoride (0.53 mL, 0.53 mmol, 1 M solution in THF) was added to a solution of di-TBS ether 15 (72.0 mg, 0.131 mmol) in THF (4 ml) under argon at 0 °C. The cooling bath was removed, and the reaction mixture was stirred for 2 h at rt. Water (5 mL) was added. The resulting mixture was extracted with ethyl ether (2 x 10 mL). The organic extracts were combined, dried with MgSO4, filtered, and concentrated under reduced pressure. The crude residue was purified via column chromatography (silica, cyclohexane/EtOAc, 80/20 to 70/30) to give the triol 16 (25 mg, 50% yield) as a colorless oil. Rf: 0.57 (EtOAc/cyclohexane: 4/6). IR (cm⁻¹): 3428, 3017, 2961, 2926, 2870, 2212, 1736, 1437, 1198, 1048, 985. ¹H NMR: (500 MHz, CDCl3): δ 6.51 (dd, J = 10.9, 15.6 Hz, 1H), 6.25-6.17 (m, 1H), 5.78-5.70 (m, 1H), 5.62-5.54 (m, 2H), 5.44-5.36 (m, 1H), 4.53-4.28 (m, 3H), 3.66 (s, 3H), 2.65-2.58 (m, 1H), 2.51-2.41 (m, 3H), 2.18-1.97 (m, 5H), 1.73-1.56 (m, 3H), 0.94 (t, J = 7.7 Hz, 3H) ppm. ¹³C NMR: (125 MHz, CDCl3): δ 171.6 (C1), 141.5 (C9), 137.0 and 136.9 (C7), 136.1 (C16), 129.9 and 129.6 (C8), 122.8 (C15), 110.9 and 110.8 (C10), 92.4 (C12), 84.1 (C11), 79.5 and 79.0 (C6), 75.9 and 75.5 (C3), 62.6 (C13), 51.8 (C19), 40.9 and 40.7 (C2), 35.7 (C14), 32.7 and 32.0 (C5), 31.9 and 31.2 (C4), 20.9 (C17), 14.4 (C18) ppm. MS: (ESI) m/z: 341.17 (M-H2O+Na)+, 301.18 (M-2H2O+1)+, 283.17 (M-3H2O+1)+, HRMS: (ESI) calculated for C₁₉H₂₆O₄Na (M-H₂O+Na)+ 341.1729, found 341.1727.

(3Z,6R,7E,9E,13S,15Z)-methyl 6,13-dihydroxyoctadeca-3,7,9,15-tetraen-11-ynoate (Diol 17 via a CSA-silyl deprotection). Catalytic CSA was added to a solution of di-TBS-ether 15 (0.116 g, 0.218 mmol) in MeOH (5 mL) at 0 °C. After stirring for 3 h at rt, the reaction was diluted with water (10 mL). The aqueous layer was extracted with EtOAc (2 x 15 mL). The combined organic phases were washed with brine (1 x 20 mL), dried over NaSO4 and evaporated, and the residue was purified by flash chromatography (silica, cyclohexane/EtOAc: 80/20 to 60/40) to afford diol 17 as a colorless oil (49 mg, 70%). Rf: 0.50 (EtOAc/cyclohexane: 4/6).
This procedure provided a 3Z/3E mixture (ca 70/30 to 85/15). The spectral data for yne-ene-diol 17 with a pure (3Z)-olefin are given below with the description of the procedure performed for the TEMPO / BAIB oxidation of primary alcohol 20.

(3Z,6R,7E,9E,13S,15Z)-methy 6,13-dihydroxyoctadeca-3,7,9,15-tetraen-11-ynoate (Diol 17 via a TEMPO/BAIB oxidation). To a stirred solution of primary alcohol 20 (14 mg, 0.027 mmol) in 
CH$_2$Cl$_2$/H$_2$O (1/1, 0.2 mL + 0.2 mL) were added BAIB (26 mg, 0.081 mmol) and TEMPO (1.2 mg, 0.0081 mmol) at rt. After stirring at the same temperature for 1 h, the reaction mixture was diluted with H$_2$O, extracted with CH$_2$Cl$_2$ (2 x 5 mL). The combined organic layers were washed with sat. Na$_2$S$_2$O$_3$ solution (1 x 10 mL) and brine (1 x 10 mL), dried over MgSO$_4$ then evaporated under reduced pressure. The crude residue was used without further purification for next step.

Catalytic CSA was added to a solution of the above-mentioned carboxylic acid (14 mg, 0.027 mmol) in MeOH (1 mL) at rt. After stirring for 1 h at rt, the reaction was diluted with sat.NaHCO$_3$ (5 mL). The aqueous layer was extracted with EtOAc (2 x 5 mL). The combined organic phases were washed with brine (1 x 10 mL), dried over MgSO$_4$ and evaporated, and the residue was purified by flash chromatography (silica, cyclohexane/EtOAc: 80/20 to 60/40) to afford the desired diol 17 as a colorless oil (4.3 mg, 46%, two steps). R$_f$: 0.50 (EtOAc/cyclohexane: 4/6). IR (cm$^{-1}$): 3397, 3017, 2961, 2928, 2873, 1736, 1437, 1333, 1167, 988. ¹H NMR: (500 MHz, CDCl$_3$): δ 6.53 (dd, J = 10.9, 15.5 Hz, 1H), 6.28 (dd, J = 10.9, 15.1 Hz, 1H), 5.79 (dd, J = 6.0, 15.1 Hz, 1H), 5.74-5.68 (m, 1H), 5.64-5.57 (m, 3H), 5.44-5.38 (m, 1H), 4.50 (br.t, J = 7.4 Hz, 1H), 4.24 (q, J = 6.0 Hz, 1H), 3.67 (s, 3H), 3.15-3.03 (m, 2H), 2.52-2.42 (m, 2H), 2.31 (t, J = 6.9 Hz, 2H), 2.16 (br.s, 1H, OH), 2.10-2.03 (m, 2H), 1.93 (br.s, 1H, OH), 0.96 (t, J = 7.6 Hz, 3H) ppm. ¹³C NMR: (125 MHz, CDCl$_3$): δ 172.6, 141.4, 138.0, 136.2, 129.5, 128.4, 124.6, 122.8, 111.0, 92.5, 84.1, 71.2, 62.7, 52.2, 35.7, 35.5, 32.9, 20.9, 14.4 ppm. MS: (ESI) m/z: 341.17 (M+Na)$^+$, 301.18 (M-H$_2$O+H)$^+$, 283.17 (M-2H$_2$O+H)$^+$.HRMS: (ESI) calculated for C$_{19}$H$_{26}$O$_4$Na (M+Na)$^+$ 341.1729, found 341.1732. HRMS: (ESI) calculated for C$_{19}$H$_{25}$O$_3$ (M-H$_2$O+H)$^+$ 301.1804, found 301.1810. HRMS: (ESI) calculated for C$_{19}$H$_{23}$O$_2$ (M-2H$_2$O+H)$^+$ 283.1696, found 283.1702.
(5R,6E,8E,12S)-2,2,3,3,14,14,15,15-octamethyl-12-((Z)-pent-2-enyl)-5-((Z)-5-(tetrahydro 2H-pyrano-2-yloxy)pent-2-enyl)-4,13-dioxa-3,14-disilahexadeca-6,8-dien-10-yn-1-ol (THP-O-protected (Z)-Wittig product 19). To a suspension of triphenyl(3-((tetrahydro-2H-pyrano-2-yl)oxy)propyl)phosphonium bromide\(^8\) (0.405 g, 0.835 mmol, prior to use, the phosphonium salt was washed with anhydrous benzene [3 x 8 mL] then dried under vacuum and over P\(_2\)O\(_5\) for 48 h at 80 °C) in anhydrous THF (9 mL) was added NaHMDS (0.35 mL, 0.7 mmol, 2M solution in hexane) at -78 °C. The temperature was raised to -10 °C. After stirring for further 30 min at the same temperature, the mixture was cooled down to -78 °C. The aldehyde (0.048 g, 0.129 mmol) was cannulated dropwise to the above prepared ylide. Then, the temperature was raised to 0 °C in 2 h. The reaction mixture was quenched with sat. NH\(_4\)Cl (10 mL) and allowed to warm to rt. Upon extraction with ethyl ether (2 x 15 mL), the combined organic layers were washed with brine (1 x 15 mL), then dried over NaSO\(_4\), filtered, and evaporated under reduced pressure. The crude residue was purified via column chromatography (silica, cyclohexane/ethyl ether, 98/02) to provide the desired Wittig product 19 (47 mg, 61% yield) as a colorless oil. \(R_f\): 0.77 (EtOAc/cyclohexane: 1/9). \(\text{IR (cm}^{-1})\): 2955, 2929, 2857, 1472, 1254, 1078, 1033, 835. \(^1\text{H NMR:}\) (300 MHz, CDCl\(_3\)): \(\delta 6.48\) (dd, \(J = 10.9, 15.5\) Hz, 1H), 6.16 (dd, \(J = 10.9, 15.2\) Hz, 1H), 5.74 (dd, \(J = 5.9, 15.2\) Hz, 1H), 5.55 (dd, \(J = 1.2, 15.6\) Hz, 1H), 5.51-5.31 (m, 4H), 4.57 (t, \(J = 2.8\) Hz, 1H), 4.45 (td, \(J = 1.6, 6.5\) Hz, 1H), 4.43 (q, \(J = 5.9\) Hz, 1H), 3.91-3.79 (m, 1H), 3.76-3.63 (m, 1H), 3.54-3.43 (m, 1H), 3.42-3.32 (m, 1H), 2.41 (t, \(J = 6.6\) Hz, 2H), 2.37-2.21 (m, 4H), 2.05 (quin, \(J = 7.2\) Hz, 2H), 1.89-1.61 (m, 2H), 1.60-1.43 (m, 4H), 0.94 (t, \(J = 7.8\) Hz, 3H), 0.88 (s, 9H), 0.87 (s, 9H), 0.11 (s, 3H), 0.09 (s, 3H), 0.02 (s, 3H), 0.00 (s, 3H) ppm. \(^{13}\text{C NMR:}\) (75 MHz, CDCl\(_3\)): \(\delta 141.2, 139.2, 134.4, 128.7, 128.0, 127.2, 124.1, 110.7, 98.9, 93.4, 83.6, 72.8, 67.1, 63.7, 62.4, 36.8, 36.5, 30.9, 28.4, 26.0 (2C), 25.6, 20.9, 19.7, 18.4 (2C), 14.4, -4.3 (2C), -4.6, -4.8 ppm. \(\text{MS:}\) (ESI) m/z: 625.41 (M+Na\(^+\)). \(\text{HRMS:}\) (ESI) calculated for C\(_{35}\)H\(_{62}\)O\(_4\)NaSi\(_2\) (M+Na\(^+\)) 625.4084, found 625.4080.

(3Z,6R,7E,9E,13S,15Z)-6,13-bis(tert-butyldimethylsilyloxy) octadeca-3,7,9,15-tetraen-11-yn-1-ol (Primary alcohol 20). To a stirred solution of THP-O-protected alcohol 19 (65 mg, 0.107 mmol) in
MeOH (3 mL) was added catalytic amount of PPTS at rt. The temperature was raised to 45 °C. After 1.5 h at 45 °C, monitoring by TLC showed that almost all starting material was consumed. It was diluted with sat. aqueous NaHCO₃ (10 mL), and the resulting aqueous phase was extracted with EtOAc (2 x 10 mL). The combined organic layers were washed with brine (1 x 10 mL), then dried over MgSO₄, filtered, and evaporated under reduced pressure. Purification by chromatography on silica gel (cyclohexane/EtOAc: 95/05) afforded the expected primary alcohol 20 (20 mg, 36%) as a colorless oil. \textbf{Rf}: 0.61 (EtOAc/Cyclohexane: 2/8). \textbf{IR} (cm⁻¹): 3364, 2956, 2929, 2857, 1472, 1253, 1078, 984, 835. \textbf{¹H NMR}: (300 MHz, CDCl₃): \(\delta\) 6.48 (dd, \(J = 10.8, 15.5\) Hz, 1H), 6.16 (dd, \(J = 10.8, 15.2\) Hz, 1H), 5.73 (dd, \(J = 6.1, 15.2\) Hz, 1H), 5.60-5.33 (m, 5H), 4.45 (td, \(J = 1.7, 6.6\) Hz, 1H), 4.20 (q, \(J = 6.1\) Hz, 1H), 3.61 (t, \(J = 6.3\) Hz, 2H), 2.41 (t, \(J = 6.7\) Hz, 2H), 2.37-2.18 (m, 4H), 2.05 (quin, \(J = 7.5\) Hz, 2H), 1.56 (br.s, 1H, OH), 0.94 (t, \(J = 7.5\) Hz, 3H), 0.88 (s, 9H), 0.87 (s, 9H), 0.11 (s, 3H), 0.09 (s, 3H), 0.02 (s, 3H), 0.00 (s, 3H) ppm. \textbf{¹³C NMR}: (75 MHz, CDCl₃): \(\delta\) 141.0, 139.0, 134.4, 128.9, 128.8, 127.6, 124.1, 110.9, 93.6, 83.5, 72.8, 63.7, 62.3, 36.8, 36.5, 31.2, 26.0 (2C), 20.9, 18.4 (2C), 14.4, -4.3 (2C), -4.6, -4.8 ppm. \textbf{MS}: (ESI) \(m/z\): 255.17 (M-2TBSOH+H)⁺, 387.27 (M-TBSOH+H)⁺, 541.35 (M+Na)⁺. \textbf{HRMS}: (ESI) calculated for C₃₀H₅₄O₃NaSi₂ (M+Na)⁺ 541.3509, found 541.3506.

The PPTS-catalyzed deprotection conditions were not selective enough. About 30 mg of corresponding triol and diols are collected and reprotected as a tri-TBS ether using TBSCl (75 mg, 0.5 mmol and imidazole (68 mg, 1.0 mmol) in CH₂Cl₂ (2 mL) at rt. Subsequently, the resulting tri-TBS ether (40 mg, 0.077 mmol) was subjected to selective deprotection using PPTS (14 mg, 0.077 mmol), providing the desired primary alcohol 20 in 63 % (20 mg) yield over 2 steps.

(3Z,6R,7E,9E,11Z,13S,15Z)-methyl 6,13-dihydroxyoctadeca-3,7,9,11,15-pentaenoate (E,E,Z-triene 21). The E,E-ynediene 17 (24 mg, 0.0754 mmol) in MeOH (5 mL) was added to a suspension of Zn(Cu/Ag) mixture (0.90 g) in water (5.0 mL). The reaction was stirred at 40 °C for 1.5 h. Acetonitrile (15 mL) was added. The reaction mixture was filtered through pad of celite, and washed with EtOAc (20 mL), dried over Na₂SO₄, and then removal of the solvents afforded a residue that was purified by column
chromatography (deactivated silica, hexane/EtOAc: 80/20 to 75/25), providing the E,E,Z-triene 21 (10.0 mg, 42% yield) as a colorless oil. \( R_f \): 0.40 (EtOAc/cyclohexane: 4/6). \textbf{IR (cm}^{-1}): 3401, 3015, 2960, 2934, 2875, 1737, 1436, 1331, 1168, 1034, 996. \textbf{\textsuperscript{1}H NMR}: (500 MHz, CDCl\textsubscript{3}): \( \delta \) 6.48 (dd, \( J = 11.8, 14.5 \) Hz, 1H), 6.34-6.15 (m, 2H), 6.06 (t, \( J = 11.2 \) Hz, 1H), 5.79-5.66 (m, 2H), 5.65-5.58 (m, 1H), 5.57-5.49 (m, 1H), 5.43 (t, \( J = 9.7 \) Hz, 1H), 5.36-5.27 (m, 1H), 4.57 (q, \( J = 7.2 \) Hz, 1H), 4.29-4.16 (m, 1H), 3.67 (s, 3H), 3.14-3.01 (m, 2H), 2.42-2.18 (m, 4H), 2.11 (br.s, 1H, OH), 2.05 (quin, \( J = 7.0 \) Hz, 2H) 1.69 (br.s, 1H, OH), 0.95 (t, \( J = 7.5 \) Hz, 3H) ppm. \textbf{\textsuperscript{13}C NMR}: (125 MHz, CDCl\textsubscript{3}): \( \delta \) 172.3, 136.4, 135.5, 133.8, 133.7, 130.5, 130.1, 128.8, 128.0, 124.4, 123.6, 71.5, 67.8, 52.2, 35.6, 35.5, 33.0, 20.9, 14.4 ppm. \textbf{MS}: (ESI) m/z: 343.19 (M+Na)\(^+\), 303.20 (M-H\textsubscript{2}O+H)\(^+\), 285.19 (M-2H\textsubscript{2}O+H)\(^+\). \textbf{HRMS}: (ESI) calculated for C\textsubscript{19}H\textsubscript{28}O\textsubscript{4}Na (M+Na)\(^+\) 343.1885, found 343.1881. \textbf{HRMS}: (ESI) calculated for C\textsubscript{19}H\textsubscript{27}O\textsubscript{3} (M-H\textsubscript{2}O+H)\(^+\) 303.1960, found 303.1957. \textbf{HRMS}: (ESI) calculated for C\textsubscript{19}H\textsubscript{25}O\textsubscript{2} (M-2H\textsubscript{2}O+H)\(^+\) 285.1855, found 285.1855.

\textbf{Other compounds}

The syntheses of \textbf{NPD1} and \textbf{dinor-NPD1 7} have been performed according to our previously\textsuperscript{6,40} reported synthesis of protectin isomers, using slighly modified procedures. Thus, the procedures for the chemical synthesis of all the precursors and targets discussed in the present manuscript, their characterization (NMR, MS, IR, UV) and their NMR spectra have been collected in the supporting information file.

\textbf{Quantitative determination of NPD1, dinor-NPD1 (7), and tetranor-NPD1 (8)}.

Quantitative determinations of \textbf{NPD1}, \textbf{dinor-NPD1 (7)}, and \textbf{tetranor-NPD1 (8)}, were carried out by RP/HPLC and multiple reaction monitoring (MRM)-mass spectrometry using the same column and solvents but shortening the gradient (%B from 65 to 100 in 6 minutes). The effluent from the HPLC was
directly interfaced into the electrospray source of a triple quadrupole (ABSciex API 4000) operated in negative ion mode. For each compound 2 different fragments arising from the molecular ion (M-H) were selected, with the most abundant used as a quantifier and the second as a qualifier ion. \(d_4\)-LTB\(_4\), a similar dihydroxylated, conjugated triene derived from arachidonic acid instead of DHA, was used as internal standard, monitoring the abundant fragmentation observed at the transition m/z 339 to 197 as a quantifier. With an RP-HPLC retention time very close to \textbf{NPD1}, \(d_4\)-LTB\(_4\) represents an excellent internal standard, sharing with \textbf{NPD1} a conjugated triene system flanked by two hydroxyl groups. Furthermore, being a stable isotope analog, \(d_4\)-LTB\(_4\) is clearly devoid of potential interferences from the corresponding endogenous compound.

Standard curves obtained always showed a correlation coefficient \(r>0.99\) (Supporting Figure S2); accuracy averaged 98.8\% throughout the different levels of the standard curve, and repeated analysis showed a CV of 7.3\% at 1 ng/ml of NPD1 and a limit of quantitation of 0.1 ng/ml, (S/N > 10).

**Human urine analysis.**

First morning urines were collected from 6 de-identified healthy donors (3 males and 3 females; 36.6±8.9 yo), that had not taken NSAIDs over the last 7 days; 1 mL aliquots were added with BHT (5 µg/mL, final concentration) and 5 ng of \(d_4\)-LTB\(_4\) as internal standard, and stored at -30°C until analysis. Separate samples were spiked with synthetic \textbf{NPD1} (50 ng) and maintained at room temperature for up to 24h, either in presence or absence of BHT, before adding internal standard and storage at -30°C. Upon thawing, samples were extracted on Strata-X (Phenomenex, Torrance, CA) solid phase cartridges and analysis of NPD1, \textbf{dinor-NPD1} and \textbf{tetranor-NPD1} was carried out by LC/MS/MS as described above.
PMN Isolation and LTB$_4$-induced Chemotaxis.

Peripheral blood human PMN were purified as previously described. In brief, PMNL were isolated by density centrifugation over Percoll ($d=1.077$) from whole blood obtained from de-identified healthy donors that had not taken NSAIDs over the last 7 days. Red blood cells were lysed by hypotonic lysis. Chemotaxis experiments were performed in a 48-well modified microchemotaxis chamber (Neuro Probe, Gaithersburg, MD, USA) using 5 µm nucleoporepolyvinylpyrrolidine-free polycarbonate filters.

Lower chambers were filled with RPMI supplemented with 0.1% bovine serum albumin (BSA) and LTB$_4$ (10 nM) as chemotactic stimulus. Negative controls were performed using the solvent of LTB$_4$ (EtOH). Upper chambers were filled with 50 µL cell suspension (2×10$^6$ cells/mL in RPMI 1640 with 0.1% BSA); NPD1, dinor-NPD1 or tetranor-NPD1 (0.01 - 10 nM) or their solvent (EtOH) were added both in the upper and lower compartment. After 90 min incubation at 37°C non-migrating cells on the upper filter surface were removed by scraping. The cells migrated to the lower side of the filter were stained with Diff-Quick stain (VWR Scientific Products, Bridgeport, NJ, USA), and densitometric analysis was performed using the Image J 1.47v software (NIH, USA). Each experiment was performed in sextuplicate and repeated with PMNL from 3-4 different donors.

**In vivo activity in acute pulmonary inflammation and FACS analysis.**

Female FVB (7–8 week old) mice were purchased from Harlan Laboratories Italy (S. Pietro al Natisone, Udine, Italy). Animals were maintained under conventional housing conditions. Prior to use, animals were acclimated for at least 5 days to the local vivarium conditions (room temperature: 20–24°C; relative humidity: 40–70%), having free access to standard rat chow and tap water. All animal experiments were carried out in agreement with the revised “Guide for the Care and Use of Laboratory Animals” and were approved by the Institutional Animal Care and Use Committee at Chiesi Farmaceutici. Intratracheal (i.tr.)
challenged with LPS was carried out using 50 µL of LPS solution (250 µg/mL in phosphate buffered solution [PBS]) containing 10 ng of tetranor-NPD1 or its solvent (EtOH, 2% final). Control animals received 50 µL of PBS with 2% EtOH. BAL collection was performed 24 hours after LPS challenge. Mice were sacrificed by an overdose of isoflurane and a midline neck incision was performed to cannulate the trachea. Lungs were washed three times with 0.6 ml of PBS as previously described. Cells were isolated by centrifugation and stained with the proper amounts of antibodies. The antibody panel was designed as follows: CD45 PE-Cy5 (BD) to first positively select all leucocytes and gate out debris, F4/80 Alexa488 (eBioscience) to gate out macrophages, Lys6G-PE (BD) and CD11b PE-Cy7 to positively identify neutrophils. BD Truecount™ Tubes were used to determine the absolute number of neutrophils in BAL. Acquisition and analysis was performed on a FACS Canto II (Becton Dickinson) using Diva 7.0 software.

ASSOCIATED CONTENT

*Supporting Information

This material is available free of charge via the Internet at http://pubs.acs.org and shows:

- Chemical procedures for the synthesis of targets 7, 8 and NPD1 and all their precursors
- \(^1\)H and \(^{13}\)C NMR spectra
- Purity: HPLC chromatograms for NPD1 and its metabolites 7 and 8
- **Figure S1.** MS/MS fragmentation spectrum of the negative ion at m/z 375 – assigned to compound 22-OH-NPD1
- **Figure S2.** LC-MS/MS with selected reaction monitoring of HepG2 cell medium for NPD1, dinor-NPD1 7, and tetranor-NPD1 8
- **Figure S3.** Calibration curves of NPD1, dinor-NPD1 (7), and tetranor-NPD1 (8) obtained using \(d_\alpha\)-LTB\(_4\) as internal standard
- SMILES molecular strings formulas (CSV)
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Author Contributions

LB, TD, AS designed the experiments, carried out the critical analysis of results, drafted and approved the manuscript; GD, LB carried out the chemical syntheses; PR, CB carried out the cell culture experiments and the mass spectrometric analysis; FS carried out the in vivo experiments in mice; AT and CB carried out the LTB₄-induced chemotaxis assays; GR performed data analysis; GD, PR, CB, AT, FS, GR, CB contributed to, and approved the manuscript.

Notes

The authors declare no competing financial interest.

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ABBREVIATIONS USED

APCI, atmospheric pressure chemical ionization; AT, aspirin-triggered; BAIB, bis(acetoxyiodo)benzene; BAL, Bronchoalveolar lavage; BHT, 2,6-Di-tert-butyl-4-methylphenol; BOR, Bestmann Ohira reaction; br s, broad signal; n-BuLi, n-butyllithium; cat., catalytic; CoA, co-enzyme A; CSA, camphorsulfonic acid; d₄-LTB₄, 5(S),12(R)-dihydroxy-6Z,8E,10E,14Z-eicosatetraenoic acid; EPA, eicosapentaenoic acid; FBS, fetal bovine serum; FVB, Friend Virus B-Type; Imid., imidazole; LPS, lipopolysaccharide; LTE₄, leukotriene E₄ – 5S-hydroxy-6R-(S-cysteinyl)-7E,9E,11Z,14Z-eicosatetraenoic acid; NaHMDS,
sodium bis(trimethylsilyl)amide; n-BuLi, n-butyllithium; [Ox.], oxidation; PMN, polymorphonuclear neutrophils; PMNL, polymorphonuclear leukocyte; PPTS, p-toluenesulfonic acid pyridine salt; Quant., quantitative; RP/HPLC, reverse phase high pressure liquid chromatography; RPMI, Roswell Park Memorial Institute medium; rt, room temperature; sat., saturated; SPM, Specialized Pro-resolving Mediators; TBDPS, tributyldiphenylsilyl; TEMPO, 2,2,6,6-tetramethyl-1-piperidinyloxy free radical; TES, triethylsilyl.

References


SYNOPSIS (Word Style “SN_Synopsis_TOC”).