Functional Metabolomics Reveals Novel Active Products in the DHA Metabolome

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Endogenous mechanisms for successful resolution of an acute inflammatory response and the local return to homeostasis are of interest because excessive inflammation underlies many human diseases. In this review, we provide an update and overview of functional metabolomics that identified a new bioactive metabolome of docosahexaenoic acid (DHA). Systematic studies revealed that DHA was converted to DHEA-derived novel bioactive products as well as aspirin-triggered forms of protectins (AT-PD1). The new oxygenated DHEA-derived products blocked PMN chemotaxis, reduced P-selectin expression and platelet-leukocyte adhesion, and showed organ protection in ischemia/reperfusion injury. These products activated cannabinoid receptor (CB2 receptor) and not CB1 receptors. The AT-PD1 reduced neutrophil (PMN) recruitment in murine peritonitis. With human cells, AT-PD1 decreased transendothelial PMN migration as well as enhanced efferocytosis of apoptotic human PMN by macrophages. The recent findings reviewed here indicate that DHEA oxidative metabolism and aspirin-triggered conversion of DHA produce potent novel molecules with anti-inflammatory and organ-protective properties, opening the DHA metabolome functional roles.

Keywords: resolvins, protectins, specialized pro-resolving mediator, DHEA, neutrophil, aspirin
FIGURE 1 | The role of chemical mediators in the acute inflammatory response. At the site of tissue injury or bacterial invasion, both exogenous and endogenous chemical mediators are liberated. Classic endogenous mediators such as prostaglandins and leukotrienes dilate vasculatures, enhance permeability of capillaries, increase blood flow, and stimulate the recruitment of neutrophils (PMNs) to form inflammatory exudate. Novel chemical mediators are produced in the evolution and resolution of the exudate that regulate tissue responses (see text for details). The black arrow denotes leukocyte traffic from venules and the dashed arrow denotes exogenous, i.e., bacterial components and chemoattractants.

FIGURE 2 | The fate of DHA in its new metabolome in vivo. Via esterification, DHA enters phospholipids and triglycerides that can circulate or reside in cell membranes. Autooxidation of DHA can produce neuroisoprostanes, the DHA form of isoprostanes. Enzymatic conversion in local inflammatory exudates can generate the specialized pro-resolving lipid mediators (SPMs), including D-series resolvins, protectins, and maresins. Each of these new families of potent local mediators carries both anti-inflammatory as well as pro-resolving actions in that they stop PMN (limit their further influx) to the site of inflammation and stimulate efferocytosis (Spite and Serhan, 2010; Serhan, 2011). Docosahexaenoic acid (DHA) is unique in this regard because it has several fates in its metabolome (Figure 2). DHA undergoes
Table 1 | Key structures of the SPM: resolvins, protectins, and maresins

<table>
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<tr>
<th>Key Structure</th>
<th>Description</th>
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<tr>
<td>RvE1</td>
<td>Resolvin E1 (5S,12R,18R-trihydroxy-eicos-6Z,8E,10E,14Z,16E-pentaenoic acid)</td>
</tr>
<tr>
<td>RvE2</td>
<td>Resolvin E2 (5S,18R-dihydroxy-eicosapentaenoic acid)</td>
</tr>
<tr>
<td>RvD1</td>
<td>Resolvin D1 (7S,8R,17S-trihydroxy-4Z,9E,11E,13Z,15E,19Z-docosahexaenoic acid)</td>
</tr>
<tr>
<td>RvD2</td>
<td>Resolvin D2 (7S,16R,17S-trihydroxydocosa-4Z,8E,10Z,12E,14E,19Z-hexaenoic acid)</td>
</tr>
<tr>
<td>RvD4</td>
<td>Resolvin D4 (4S,5,17S-trihydroxydocosa-6E,8E,10Z,13Z,15E,19Z-hexaenoic acid)</td>
</tr>
<tr>
<td>RvD5</td>
<td>Resolvin D5 (7S,17S-dihydroxy-docosa-4Z,8E,10Z,13Z,15E,19Z-hexaenoic acid)</td>
</tr>
<tr>
<td>MaR1</td>
<td>Maresin 1 (7,14-dihydroxy-docosa-4Z,8,10,12,16Z,19Z-hexaenoic acid)</td>
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Complete structures of the key SPM are reported and total organic synthesis was recently reviewed in Serhan and Petasis (2011).

Because of the limited quantities of endogenous 17-HDHEA produced in brain tissue, further studies and in vitro enzymatic preparation were carried out by incubating DHEA with 15-LOX followed by reduction with NaBH₄. Endogenous 17-HDHEA and the enzymatically prepared compound in vitro gave essentially the same LC retention times and tandem mass fragmentations. To assess their production by human and mouse tissues, DHEA was also incubated with isolated human PMN or whole mouse brain because DHEA is enriched in this tissue. LC–MS–MS-based targeted lipidomics indicated the production of a novel series of oxygenated products (see Figure 4).

**DECODING**

As second approach in parallel to structure elucidation, screening of HPLC-isolated DHEA metabolites obtained from mouse brain was carried out utilizing a microfluidic chamber (Irimia et al., 2006; Kasuga et al., 2008). After a gradient of IL-8 was introduced to the main channel of the device, P-selectin tethered leukocytes rapidly migrated in the IL-8 gradient at an average rate of 2.3 μm/min. Next, a mixture of isolated metabolites was infused into the main channel. Human PMN chemotaxis is dramatically reduced upon the addition of the brain metabolite mixture. For example, the average human PMN chemotaxis velocity dropped from 2.3 to ~ 0.7 μm/min (Yang et al., 2011). These results indicated that the brain fraction of metabolites contained products that stopped PMN chemotaxis. Results from this screening uncovered that at least one bioactive product was present among the mixture of DHEA metabolites. Hence, we next pursued the metabolic fates of DHEA and 17-HpDHEA/17-HDHEA identified in mouse brain (Yang et al., 2011) using LC/UV/MS/MS-based lipidomics. As with 17-HDHEA, acetate adducts of potential DHEA-derived novel metabolites [M + CH₃COOH-H] were targeted for tandem mass analysis. From these studies we identified the presence and production of novel products in the DHEA metabolome (Figure 4).

Human PMN incubated with either DHEA or 17-HpDHEA led to the generation of 17-HDHEA, 4,17-diHDHEA, 10,17-diHDHEA, and 15-HEDPEA (see Figure 4 and Yang et al., 2011). Human hemoglobin, which can be liberated upon tissue damage (Kumar et al., 2005), was next incubated with 17-HpDHEA. This gave 13-HEDPEA and 15-HEDPEA as prominent products, as well as 17-HDHEA (see Figure 4). Mouse brain homogenates with DHEA also produced 17-HDHEA and 4,17-diHDHEA as major products with smaller amounts of 7,17-diHDHEA, 10,17-diHDHEA, and 15-HEDPEA. The adduct parent ion, analyte parent ion and the ions resulting from neutral loss are m/z 462 = [M + CH₃COOH-H], 402 = [M-H], 384 = [M-H - H₂O], 366 = [M-H - 2H₂O], which are common signature ions for all dihydroxy-containing DHEA products. The ions m/z 333, 315 = [333 – H₂O], 304, 286 = [304-H₂O] were assigned as diagnostic ions for fragmentations at position 17. Fragmentations at position 4 can lead to m/z 144, 257 and 239 = [257-H₂O]. Its UV spectrum displayed characteristic maximum absorbance at 238 nm, which was consistent with the presence of two separated conjugated diene structures in this compound. Diagnostic ions m/z 304, 286 = [304-H₂O], 184, 156, corresponded to the fragmentations at positions 7 and 17 of 7,17-diHDHEA respectively.

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Shinohara et al. Functional metabolomics
Shinohara et al. Functional metabolomics

FIGURE 3 | DHA bioactive metabolome. DHA is the precursor to D-series resolvin (RvD), aspirin-triggered resolvin D (AT-RvD), maresins (MaR), protectin D1 (PD1), and aspirin-triggered protectin D1 (AT-PD1). Another metabolic fate of DHA is conversion to docosahexaenoylethanolamide (DHEA). See text for details on the recently identified novel series of bioactive oxygenated DHEA-derived products including 15-HEDPEA and 10,17S-diHDHEA.

(Yang et al., 2011). The UV spectrum of the compound displayed maximum absorbance, $\lambda_{\text{max}}$, at 246 nm (Tjonahen et al., 2006), consistent with the presence of two diene structures separated by a methylene group. For 10,17-diHDHEA, $m/z$ 333, 315 = [333-H$_2$O], 304, 286 = [304-H$_2$O], 196 came from fragmentations at positions 10 and 17 (Yang et al., 2011). The presence of a conjugated triene structure in 10,17-diHDHEA was confirmed by the characteristic UV spectrum with $\lambda_{\text{max}}$ at 270 nm. Tandem mass spectrum of 13-HEDPEA (Yang et al., 2011) gave signature fragmentation ions $m/z$ 320, 304, 286 = [304-H$_2$O], 236. GC/MS was also utilized for additional structural analysis with 13-HEDPEA and 15-HEDPEA that helped to confirm the tandem MS assignments.

To determine concentrations, as well as to further confirm structures, HPLC-isolated 13-HEDPEA and 15-HEDPEA were characterized using proton NMR ($^1$H NMR). In the chemical shift assignments (Yang et al., 2011) for 15-HEDPEA, the proton at position 15 (H-15) displayed two distinct chemical shifts. Because of limited amounts of materials and the lack of informative UV chromophores present in these compounds, NMR spectroscopy was also used for quantitation using 17-HDHA as an internal standard with known concentrations. Each NMR quantitated compound was then used for HPLC calibration and quantitation (Yang et al., 2011).

Each of these HPLC-isolated dioxygenated DHEA products were screened for direct PMN actions using microfluidic chambers and freshly isolated PMN (Yang et al., 2011). Infusion of isolated 15-HEDPEA at 10 nM to the main channel stimulated changes in morphology and chemotaxis of human PMN in the IL-8 gradient and stopped further PMN migration (Figure 4). For direct comparison, PMN chemotaxis velocity did not change with time with the IL-8 gradient. Also, at 10 nM, 4,17-diHDHEA, 7,17-diHDHEA, or 10,17-diHDHEA did not significantly regulate chemotaxis, whereas at higher concentrations, e.g., 10 $\mu$M, 10,17-diHDHEA rapidly stopped PMN. Together, these results indicated that 15-HEDPEA is the most potent of this series in regulating human PMN shape change and motility.

CANNABINOID RECEPTORS AND NOVEL DHEA PRODUCTS
Because AEA exerts a wide range of bioactions via activating cannabinoid receptor(s) (Devane et al., 1992; Felder et al., 1993), we therefore tested whether DHEA, 10,17-diHDHEA or 15-HEDPEA also activated CB receptors. For these experiments, we used recombinant human CB receptors over-expressed in a
DHEA is first converted to 17-HpDHEA by 15-LOX. Next, 17-HpDHEA is partially reduced to the oxide radical, which reacts with the vicinal double bond at the 16-position to yield the 16(17)-epoxide radical (see text and Yang et al., 2011). Non- or low stereospecific addition of oxygen to the intermediate leads to formation of two types of peroxide radical diastereomers; their further reduction produces 13-HEDPEA and 15-HEDPEA. Via a LOX-related mechanism, 17-HpDHEA is converted to 10, 17S-diHDHEA or directly reduced to 17S-HDHEA, 10, 17S-diHDHEA, and 15-HEDPEA block PMN chemotaxis, reduce P-selectin expression and aggregation of platelets and leukocytes, and show organ protection in ischemia/reperfusion injury. Although these new products directly act on recombinant CB2 receptors in vitro, the activation of CB2 in vivo and/or additional receptors in vivo by 10, 17S-diHDHEA and 15-HEDPEA remains of interest.
beta-arrestin system. HPLC purified AEA was used for direct comparison as a known receptor agonist of CB1 (Janero and Makriyannis, 2007; Vemuri et al., 2008). In these experiments, activation of CB2 by AEA gave EC50 ~ 1.1 × 10^−10 M and DHEA 9.8 × 10^−9 M. For comparison, EC50 for metabolically oxygenated products, 10,17-diHDHEA and 15-HEDPEA, were 3.9 × 10^−10 M and 1.0 × 10^−10 M (Yang et al., 2011). These results demonstrated that the novel enzymatic oxidation products from DHEA are activators of CB2 receptors, and that 10,17-diHDHEA and 15-HEDPEA also activated CB1 receptors but required much higher concentrations. In this system, 15(S)-HETE ethanolamide (oxygenated product of AEA) did not stimulate CB2 receptors in this dose range. CB2 receptor-ligand interactions were confirmed with dose response of CB2 specific antagonist AM630. When incubated with GPCR CB2 over-expressed cells, AM630 inhibited activation stimulated with 15-HEDPEA (10 nM) and AEA (10 nM). AM630 also inhibited GPCR CB2 interaction with 10,17-diHDHEA at higher concentration (Yang et al., 2011).

**NOVEL DHEA PRODUCTS REDUCED PLATELET-LEUKOCYTE AGGREGATE**

Platelet-leukocyte interactions play important roles in hemostasis, thrombosis, and inflammation (for a recent review see van Gils et al., 2009 and references within). At concentrations as low as 10 pM, 10,17S-diHDHEA or 15-HEDPEA decreased PAF (100 nM) stimulated platelet-monocyte aggregate formation in human whole blood by ~30% (Yang et al., 2011). The inhibitory action of 10,17-diHDHEA displayed a bell-shaped dose response and reached maximum reduction at ~40% with 100 pM. The formation of PMN-platelet aggregates with PAF (100 nM) was also inhibited by 10,17-diHDHEA at concentrations as low as 10 pM, as was the surface expression of P-selectin on platelets in human whole blood (Figure 4). By comparison, the precursor DHEA was not active in this dose range.

**NOVEL DHEA PRODUCTS ARE ORGAN PROTECTIVE IN ISCHEMIA/REPERFUSION INJURY**

15-HEDPEA displayed potent bioactions with human PMN at single-cell level (Figure 4 and Yang et al., 2011) and in human whole blood; hence, we questioned whether it had protective actions in vivo in murine hind limb ischemia (1 h) and second organ reperfusion (2 h) injury (Qiu et al., 2000). Indeed, following reperfusion, 15-HEDPEA significantly reduced lung PMN accumulation in mice and associated lung injury at 1 μg/mouse (Figure 4 and Yang et al., 2011; ~50% reduction directly compared with vehicle).

Lipidomic investigation of DHEA functional metabolome uncovered a series of novel oxygenated products that (1) are potent CB2 agonists, (2) regulate single-cell PMN chemotactic responses, (3) modulate platelet-leukocyte interaction in whole blood, and (4) are organ protective. In view of the role of lipid mediators in inflammation and its resolution as well as homeostasis (Serhan et al., 2008), the present new DHEA metabolome documented herein may serve as a counter-regulatory system in neural tissues and those rich in DHEA as well as in administration of DHA (Calder, 2010) to regulate leukocyte-mediated tissue damage. The link between CB2 receptor activation in vitro and in vivo actions of these novel products remains to be established. Also, it is possible that additional receptors in vivo may be functionally regulated by these new DHEA products.

**ASPIRIN AND DHA: UNCOVERING THE AT-(NPD1/PD1) PATHWAY**

Aspirin is well appreciated for its ability to inhibit COX-1 and inactivate this enzyme. On the other hand, the catalytic region of COX-2 is larger than that of COX-1, and when it is acetylated by aspirin, acetylated COX-2 remains active, producing lipoxygenase-like products, demonstrated in vivo in humans (Chiang et al., 2004; Chiang and Serhan, 2004). Aspirin is unique in that it can jump-start resolution by triggering production of E-series and D-series resolvins and AT-lipoxins (Serhan et al., 2000, 2002; Schwab et al., 2007; Morris et al., 2009). The previously unrecognized mediators and mechanisms involve the biosynthesis of AT lipid mediators. With Dr. Bazan (LSU) and Dr. Petasis (USC), we established DHA conversion to the AT-NPD1 pathway (Serhan et al., 2011).

To determine the complete stereochemical assignment and bioactions of AT-(NPD1/PD1), we directly compared the physical and biological properties of DHA-derived AT-(NPD1/PD1) and related 10,17 dihydroxy-docosatriene stereoisomers produced by leukocytes to those prepared in stereochemically pure form by total organic synthesis (Serhan et al., 2011). Since AT-(NPD1/PD1) was identified in resolving murine exudates treated with aspirin (Serhan et al., 2002), biologic AT-(NPD1/PD1) was obtained from resolving murine exudates in vivo. To this end, peritonitis was initiated via intra-peritoneal (i.p.) administration of zymosan A (1 mg/mouse), and exudates were harvested 24 h post injection (i.e., within the resolution phase; Bannenberg et al., 2005). Exudates were subject to solid phase extraction and analyzed using LC–UV–MS–MS-based mediator lipidomics. AT-(NPD1/PD1) was also isolated and identified from activated human leukocytes, namely, aspirin-treated human PMN. Collectively, by matching biologic materials from both human and murine exudate AT-(NPD1/PD1) with synthetic candidates, these results established the complete stereochecmistry of endogenous AT-(NPD1/PD1) as 10R,17R-dihydroxydocosa-4Z, 7Z, 11E, 13E, 15Z, 19Z-hexaenoic acid (Serhan et al., 2011).

**ANT-INFLAMMATORY ACTIONS OF AT-NPD1/PD1**

Earlier results indicated that NPD1/PD1 exerted potent anti-inflammatory actions regulating leukocyte trafficking in murine systems (Serhan et al., 2002; Hong et al., 2003). We compared the bioactions of NPD1/PD1 to those of AT-(NPD1/PD1) carried out in parallel with the physical matching experiments (Serhan et al., 2011). Synthetic AT-(NPD1/PD1) limited PMN infiltration into the peritoneum in TNFα-stimulated peritonitis (Figure 5). Both NPD1/PD1 (0.1–10 ng) and AT-(NPD1/PD1; 0.01–10.0 ng) proved to be significant regulators of TNFα-stimulated leukocyte infiltration into the peritoneum. AT-(NPD1/PD1) reduced total leukocyte population of the exudate including PMN, monocyte and lymphocyte infiltrates, reaching a maximal reduction at 1 ng/mouse by as much as 50.4 ± 8.8%. The PMN population was also reduced with AT-(NPD1/PD1) reaching a maximal PMN reduction at 1 ng/mouse (62.2 ± 7.8%; Serhan et al., 2011). The flow cytometry results obtained with murine exudates indicated
FIGURE 5 | Proposed biosynthesis scheme for protectin D1 (PD1) and aspirin-triggered protectin D1 (AT-PD1). DHA is first converted to 17S-HpDHA intermediate by 15-LOX. Then PD1 is generated through 16S, 17S-epoxide. In the aspirin-triggered pathway, 17R-HpDHA is generated from DHA via acetylated-COX, and converted to AT-PD1 via a 16R, 17R-containing epoxide intermediate. Both PD1 and AT-PD1 reduce leukocytes and PMN infiltration in peritonitis stimulated by either TNF-α or zymosan A. PD1 and AT-PD1 each enhance human macrophage efferocytosis of apoptotic human PMNs and limit human PMN transmigration across human endothelial cells, the defining pro-resolving bioactions.

a reduction in the Ly6G+CD11b+ population as compared to TNFα alone. Of note, the Δ15-trans-AT-(NPD1/PD1) isomer did not reduce either the total exudate leukocyte population or PMN infiltration. Since NPD1/PD1 limited PMN infiltration in zymosan A-initiated peritonitis (Serhan et al., 2006), we determined whether AT-(NPD1/PD1) also reduced PMN infiltration in zymosan A-stimulated peritonitis. AT-(NPD1/PD1; 0.1–100.0 ng) significantly reduced total leukocytes, as well as PMN infiltration, reaching a maximal reduction of 47.8 ± 10.0% and 49.1 ± 11.9%. In comparison to AT-(NPD1/PD1; 1.0 and 10.0 ng), equal doses of either the precursor DHA or Δ15-trans-AT-NPD1/PD1 did not reduce either total leukocyte infiltration or PMN
infiltration (Serhan et al., 2011). These findings indicated that AT-(NPD1/PD1) regulates inflammatory responses induced by the pro-inflammatory cytokine TNFα and the TLR ligand, zymosan A.

**PMN-TRANSENDOTHELIAL MIGRATION AND ENHANCED EFFEROCYTOSIS**

PMN-transendothelial migration is the first committed step of leukocytes in acute inflammation (Kumar et al., 2005). AT-(NPD1/PD1) and NPD1/PD1 (0.1–10.0 nM) reduced (∼30 and ∼50%) PMN-transendothelial migration stimulated by LTB4. Equal concentrations of the Δ13-trans isomer of AT-(NPD1/PD1) where the conjugated triene portion of the molecule was in the trans rather than cis configuration did not reduce PMN-transendothelial migration. Again, the precursor DHA (Kasuga et al., 2008) did not reduce LTB4-stimulated PMN-transendothelial migration. To corroborate these, we also used an electric cell-substrate impedance sensing system (ECIS) that sensitively quantitates cellular responses in two cell systems by real-time monitoring of barrier impedance (Tsikitis et al., 2004). Both AT-(NPD1/PD1) and NPD1/PD1 (1nM) decreased LTB4-stimulated PMN-transendothelial migration, and AT-(NPD1/PD1) also enhanced the uptake of apoptotic human PMN by human macrophages at concentrations as low as 0.1 nM, as did NPD1/PD1 when compared directly. The response was bell-shaped and consistent with the dose response relationship observed for effecrocytosis and pro-resolving lipid mediators such as RvE1 (Hong et al., 2008).

**SOME CONSIDERATIONS FOR THE FUTURE**

In summation, the novel products and pathway metabolome reviewed here might have protective and anti-inflammatory actions relevant in humans. This important level of evidence is of interest and remains to be established in human trials. For example, since RvE1 has a direct protective effect on cardiomyocytes against ischemia-reperfusion injury limiting infarct size when administered intravenously just before reperfusion (Keyes et al., 2010), the novel bioactive DHEA products discussed in this review may carry potential clinical applications. These new DHEA products (Figure 4) and/or related structures might be useful in the treatment of acute myocardial infarction, because platelet and neutrophil interactions have a key regulatory role in the site of inflammation (Marcus et al., 1982; Phillipson and Kubes, 2011). Along these lines, Pillai et al. (2012) from this laboratory recently reported on the time course and identification of inflammatory mediators (both cytokine and lipid mediators) after abdominal aortic aneurysm (AAA) surgery. Chemical mediator profiles from patients that had undergone AAA in that study assembled into two groups. The temporal profiles for local chemical mediators from these patients were either those with a pro-inflammatory profile of local mediators or a second group with a potential resolving profile of local mediators. These recent reports and results reviewed herein suggest that the temporal biosynthesis of local chemical mediators following inflammation or tissue injury can have an acute local impact as well as long-term effects following surgical intervention and tissue remodeling. Hence, knowledge of these new mediators and pathways as well as their potential roles may serve as a basis for new and more effective therapeutics that could improve the outcome of diseases, surgery and post-operative events.

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**REFERENCES**


Shinohara et al. Functional metabolomics


Conflict of Interest Statement: The authors declare that this review was written in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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