Prostacyclin and Interaction with Diacylglycerol Lipase and CDP Diacylglycerol; Possibility of De Novo Synthesis of Prostacyclin or Related Congeners by Novel Mechanisms

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Abstract: Prostacyclin is a strong cardioprotective hormone released by the endothelium of the blood vessels. Prostacyclin is present in equilibrium with several vasoactive agents in cardiovascular system. In recent years, prostacyclin (PGI2) has also been shown to enhance differentiation and inhibit proliferation in vascular smooth muscle cells. In addition to these well-described homeostatic roles within the cardiovascular system, prostacyclin (PGI2) also plays an important role as an inflammatory mediator. In this review, the focus on the contribution of prostacyclin (PGI2) as both a patho-physiological mediator in three major inflammatory-mediated disease processes, namely rheumatoid arthritis, where it promotes disease progression, along with pulmonary vascular disease and atherosclerosis, where it inhibits disease progression. On the other hand, CDP-DAG synthases (CDS) are enzymes that catalyze the conversion of phosphatidic acid (PA) to CDP-diacylglycerol (CDP-DAG). Both PA and CDP-DAG serve critical roles in cellular functions. This article reviews the possibility of interaction with CDP diacylglycerol and it appears that de novo synthesis of PGI2 or its congeners occurs in specialized cells under patho-physiological conditions.

Key words: Prostacyclin, diacylglycerol, enzyme, phosphatidic acid, vascular, inflammation

I. Introduction
Salvador Moncada showed in his exemplary work the importance of prostacyclin as an important autacoid and with vasodilatory properties and anti-platelet agent (1). Prostacyclin (PGI2) is a potent vasorelaxant and an inhibitor of platelet aggregation. Prostacyclin is synthesized from free arachidonic acid, and is released from membrane phospholipids, which is converted into prostaglandin endoperoxide (PGH2) by prostaglandin synthases (PGHS). PGH2 is subsequently metabolized to prostacyclin by the enzyme prostacyclin synthase. Prostaglandin I2 (PGI2) is the major metabolite of the cyclooxygenase pathway, and epoxyeicosatrienoic acids (EETs) are the products of the cytochrome P-450 epoxygenase pathway (2-4). PGI2 relaxes vascular smooth muscle by increasing cAMP and opening ATP-sensitive potassium (K+) channels (5). EETs represent endothelium-derived hyperpolarizing factors. On the hand, the roles of CDS1 and CDS2 have primarily been studied in PI synthesis. Many of the cellular functions attributed to CDS enzymes are believed to result from their role in generating the precursor for phosphatidylinositol 4,5-bisphosphate (PIP2), a potent signaling molecule (6). For example, photo-transduction signaling in vertebrate and invertebrate systems is believed to proceed, at least partly, via phosphoinositide signaling. Although PLA2 activation is the most direct route for arachidonic acid release in response to various stimuli, alternative pathways exist in some cells. Arachidonic acid can be released from diacylglycerol (DAG), generated via phospholipase C (PLC) activation, by the enzyme DAG lipase (7). Previous studies have shown that fibroblasts from cancer mass produce PGI2 but fibroblasts from adjacent normal tissues do not. However, how cancer associated fibroblasts (CAFs) are activated and if the activated CAFs can promote angiogenesis by generating VEGF itself are poorly understood. Recent experimental studies suggest the modulation of the gene expression in human fibroblasts under hypoxic condition using GeneChip analysis, and found that the expression of prostacyclin synthase (PGIS) was upregulated. PGIS, a membrane-bound heme protein with spectral characteristics of cytochrome p450 (CYP), is also an enzyme which catalyzes the conversion of prostaglandin H2 (PGH2) to form PGI2. PGIS is localized to the microsomal fractions of platelets, vascular endothelial cells, and vascular smooth muscle cells (8-9). Therefore this article evaluates the role of PGI2 in inflammation and interaction with CDP diacylglycerol and possibility of prostacyclin or related congeners synthesized by de-novo pathways.

PGI2 and signal transduction: PGI2 is unstable at physiological pH and, thus, has a very short half-life in vivo (<2 min), rapidly transforming into the inactive hydration product 6-keto-prostaglandin F1α or 6-oxo-PGF1 alpha (10-11). The actions of PGI2 are mediated through a seven-transmembrane-spanning G-protein coupled receptor (GPCR), referred to as the IP receptor (IUPHAR nomenclature) as shown in Fig.1. The IP receptor is a Class A rhodopsin-like GPCR that couples pre-dominantly to the Gs subunit of the heterotrimetric G-protein and mediates intracellular signaling via adenyl cyclase activation and cyclic AMP production (12). Des –
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aspartate-angiotensin 1 (DAA-1) peptide derivative has also been shown to release PGI2. The release of PGE2 and PGI2 via the angiotensin AT1 receptor and COX-1 is a novel specific action of DAA-1 and is likely responsible for its beneficial effects seen in earlier studies. This specific action is definable as a biased agonism of the angiotensin AT1 receptor, which identifies DAA-1 as a novel biased agonist and potential therapeutic that is able to produce specific prostaglandins at nanomolar concentration, thus throwing light on possible novel mechanism for prostaglandins as well (13). Animal studies have also shown that PGI2 may also signal through alternate Gq– and Gi-related pathways (14), as well as nuclear receptor-mediated pathways, such as the peroxisome proliferator-activated receptor gamma (PPARγ) pathway (15).

**PGI2 and postulated role in Inflammation:** The seminal work by Vane (16) demonstrating the inhibition of prostaglandin biosynthesis as the mechanism of action for aspirin (acetylsalicylic acid) and other aspirin-like drugs first highlighted the importance of the prostaglandin family of molecules, and set the stage for the development of many pharmacologic agents, such as traditional, non-selective non-steroidal anti-inflammatory drugs (NSAIDs) and the newer selective COX-2 inhibitors. Further work conducted by Davies et al. (17) pinpointed particular prostaglandins, principally prostaglandin E2 (PGE2) and prostacyclin (PGI2), in the mediation of vascular permeability associated with the hyperemia and edema seen with acute inflammation. Murata et al. (18) demonstrated the involvement of prostacyclin (PGI2)-mediated inflammatory swelling in vivo, using prostacyclin receptor deficient (IP−/−) mice. In these critical experiments, it was shown that mice lacking the prostacyclin receptor had a reduced inflammatory response, as measured by percent change in vascular permeability using a carrageenan-induced paw-edema model.

**CDP diacylglycerol synthases:** Only two CDP diacylglycerol synthases (CDS) isoforms in mammals have been identified and characterized. Both these isoforms are believed to be localized to the endoplasmic reticulum (ER). It was believed that CDS1 was present in mitochondria for synthesizing cardiolipin. CDS1 and CDS2 are expressed in a variety of tissues. In mice, CDS1 is found in adult brain, eye, smooth muscle, and testis (19). In the eyes, CDS1 is strongly expressed in the photoreceptor layer of adult retinas, which could suggest a role for CDS1 in phototransduction. CDS2 has a broad expression pattern and was found in virtually every tissue, however, some discrepancies exist in the tissue localization of CDS2 (20). The de novo synthesis of phosphoinositide (PI) involves only the endoplasmic reticulum and generates mainly saturated and monounsaturated acyl chains. The PI cycle is a cyclical pathway that involves the breakdown and regeneration of PIP2. The PI cycle involves both the endoplasmic reticulum and plasma membrane and results in the enrichment of 1-stearoyl-2-arachidonyl species. Both pathways involve common features, one of which is the conversion of PA species to CDP-DAG by CDS enzymes. It appears likely that CDS2 would be involved in the PI cycle (19). The acyl chain selectivity of CDS is similar to that of DGKε, which was shown to be required for the arachidonoyl enrichment of PI species. CDS2 could play a similar yet important role in the enrichment of PI with an arachidonoyl chain. CDP-DAG produced by CDS2 can be used only for the synthesis of phospholipids (21-22).

**Table 1:** Effect of DAG lipase inhibitor RHC 80267 and IL-1 beta on 6-oxo-PGF1 alpha levels in human lymphocyte cells (n=5)

<table>
<thead>
<tr>
<th>No.</th>
<th>Pretreatment</th>
<th>Treatment</th>
<th>6-oxo-PGF1 alpha (pg/well)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1)</td>
<td>Saline</td>
<td>Saline</td>
<td>313 ± 35.11</td>
</tr>
<tr>
<td>2)</td>
<td>Saline</td>
<td>IL-1beta (200U/ml)</td>
<td>385 ± 30.22</td>
</tr>
<tr>
<td>3)</td>
<td>Saline</td>
<td>RHC80267 (20µM/ml)</td>
<td>165 ± 23.72</td>
</tr>
<tr>
<td>4)</td>
<td>IL1-beta</td>
<td>RHC 80267</td>
<td>225 ± 27.97</td>
</tr>
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</table>

**Figure 1:** Hypoxia and prostacyclin mediated signaling pathway

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II. Discussion

Table I shows the effects of IL1 beta and DAG lipase inhibition by RHC 80267 on 6-oxo-PGF1 alpha levels as measured by radioimmunoassay. It appears that in lymphocyte cells the RHC 80267 either directly inhibits the prostacyclin synthesis or its congeners or otherwise the accumulation of diacylglycerol causes the inhibition of prostacyclin or its putative congeners via some unspecified mechanisms. On the other hand interleukin 1 beta was able to increase the levels of PG12 and this was attenuated by the known DAG lipase inhibitor RHC 80267. The work done by D’Souza et al is suggestive of the importance of CDS enzyme and apparently there is more research work required in this area to confirm the exact role played by diacylglycerol and related pathways and the synthesis of prostacyclin like congeners (23-24). Thus it is still an area where more research work needs to be carried out and the exact phenomenon elucidated.

III. Conclusions

Prostacyclin is an important prostaglandin which has important vasodilatory and anti-platelet activity and roles beyond the cardiovascular system. Our existing knowledge of PGI2, as both a physiological–pathophysiological mediator and therapeutic agent, in a host of inflammatory-related diseases, is growing rapidly. As demonstrated by several studies, PG12 has been shown to play protective roles in atherosogenesis – relating to CAD, MI, stroke, and other cardiovascular abnormalities. It has also been shown to be involved in certain fibro-proliferative and pulmonary vascular diseases, such as idiopathic pulmonary fibrosis and pulmonary hypertension, where it serves as both a protective factor and first-line pharmacotherapy. It appears from some recent studies that prostacyclin or its congeners may be synthesized by novel de-novo pathways. Further study of this important prostaglandin, in both the realms of basic science and clinical medicine, is required and will undoubtedly lead to new insights into inflammatory disorders and pharmacological treatments.

References

