Pharmacological Inhibition of Phospholipase A2: Results from Phase 3 Clinical Trials with Darapladib and Varespladib in Patients with Cardiovascular Disease

Nicola Ferri1*, Chiara Ricci1 and Alberto Corsini1,2
1Dipartimento di Scienze Farmacologiche e Biomolecolari, Università degli Studi di Milano, Italy
2IRCCS, Multimedia, Milan Italy

Abstract
The hydrolysis of the ester bond of glycerophospholipids is catalyzed by the family of enzymes Phospholipase A2 (PLA2), that leads to a release of free fatty acids and lysophospholipids, including the arachidonic acid, the precursor of the eicosanoids and the inflammatory cascades. The mass and the enzymatic activity of PLA2 have been positively correlated with the incidence of cardiovascular diseases in epidemiological and genetic studies. In particular, several experimental evidences have shown that PLA2, identified in the atherosclerotic plaque, are directly involved in the proatherogenic inflammatory response. From these evidences, PLA2 have become a potential pharmacological target of considerable interest and two different PLA2 inhibitors have been developed: varespladib, a reversible sPLA2 inhibitor, and darapladib, a selective Lp-PLA2 inhibitor. Both these two small molecules have been tested both on animal models, where they have shown anti-atherosclerotic properties, and in phase 2 clinical trials, where they have demonstrated positive effects on atherosclerotic plaque composition. Unfortunately, the following three phase 3 trials, which have been recently published, did not shown any additional protective action of PLA2 inhibitors neither in co-administration with statins and antiplatelet drugs, nor in coronary revascularization. In the first one, the VISTA-16 study, varespladib has been administrated to patients with acute coronary syndrome, in the second and third one, the Stability and the SOLID-TIMI 52 studies, darapladib has been administrated to patients with stable coronary heart disease and acute coronary syndrome, respectively. The present article is focused on the enzymatic properties and on the involvement of sPLA2 and Lp-PLA2 in atherogenesis, with particular attention on the results of experimental and clinical studies with both varespladib and darapladib.

Keywords: Darapladib; Cardiovascular diseases; Atherosclerosis; Varespladib

Introduction
In the recent years, it has been documented a significant reduction in morbidity and mortality from cardiovascular causes. Nevertheless, it is evident that even in patients treated aggressively with currently available drugs, the rate of cardiovascular events still remains high. These conditions highlight the need to develop new strategies for the treatment of cardiovascular disease. In the mid 90, several evidences have support the hypothesis for the key role of vascular inflammation in atherogenesis, as documented by the involvement of pro-inflammatory molecules, such as C-reactive protein (CRP), interleukin-1 (IL-1), p38 MAPK and phospholipase A2, both soluble (sPLA2) or associated with lipoproteins (Lp-PLA2). The role of inflammatory response in atherogenesis has also been confirmed by the evidence of the so called "pleiotropic" effects of statins (the most potent antiatherosclerotic agents currently available in clinic) that could affect the vascular response to injury and reduces the inflammatory marker CRP [1-8]. The Lp-PLA2, in particular, has been considered to be a potential pharmacological target for the development of new drugs with anti-atherosclerotic activity [9].

Role of sPLA2 and Lp-PLA2 in the Atherosclerosis
sPLA2
The PLA2, represent a class of enzymes that hydrolyze the sn-2 ester bond of glycerophospholipids leading to the formation of free fatty acids and lysophospholipids, such as arachidonic acid, the precursor of the eicosanoids. The soluble PLA2 are divided into 10 groups, which include 13 different isoforms; in particular, the sPLA2-I, the sPLA2-III, the sPLA2-V and the sPLA2-X are involved in atherogenesis (Table 1) [10,11]. The four isoforms show a different capacity to hydrolyze the phosphatidylcholine (PC) and phosphatidylethanolamine (PE) and the sPLA2-V has a unique role in hydrolyzing phospholipids present in human lipoproteins [12-14]. All these four isoforms of soluble PLA2 are present in the atherosclerotic plaques [15,16], although with a different distribution. The sPLA2-V is mainly expressed in smooth muscle cells while the other (IIA, III, and X) are also present in macrophages [13]. Regarding the role of sPLA2 in atherogenesis, it is generally accepted, that these enzymes are capable to modify the LDL and increase the ability to bind to proteoglycans of the extracellular matrix present in the vessel wall facilitating their aggregation and oxidation. The enzymatic activity of both sPLA2 and Lp-PLA2 leads to the formation of bioactive fatty acids (such as arachidonic acid) and lysophosphatidylcholine (lyso-PC) capable to promote cell activation and the production of pro-inflammatory cytokines. The sPLA2 also promotes the formation of macrophage-derived foam cells by modifying the lipoprotein particles.

Lp-PLA2
Differently from the sPLA2, there is only one isoform of Lp-PLA2.

*Corresponding author: Nicola Ferri, Dipartimento di Scienze Farmacologiche e Biomolecolari, Università degli Studi di Milano, Italy, Via Balzaretti 9, 20133 Milano, Tel: 0250318322; E-mail: nicola.ferri@unimi.it

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Pharmacological properties

<table>
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<th>Structural features</th>
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<tr>
<td>sPLA₂-V</td>
<td>13.8</td>
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<tr>
<td>sPLA₂-X</td>
<td>13.6</td>
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<tr>
<td>Lp-PLA₂</td>
<td>45</td>
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</tbody>
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PE=Phosphatidylethanolamine; PC=Phosphatidylcholine; sPLA₂ and Lp-PLA₂ have different molecular masses and different active sites: a His/Asp dyad and a Ser/Asp/His triad, respectively.

Table 1: Structural and functional features of sPLA₂ and Lp-PLA₂

Darapladib Pharmacological properties

- Inhibition of Lp-PLA₂; IC₅₀=0.25 nM [28]
- Reduction of Lp-PLA₂ activity in experimental and human atherosclerotic lesions [34]
- Reduction of Lp-PLA₂ activity in carotid plaques [35]
- Inhibition of the progression of the necrotic core volume of coronary plaques in the IBIS-2 trial conducted in patients with cardiovascular disease [33]

Experimental models

- Reduction of atherosclerotic plaque area and total cholesterol levels in mouse models of atherosclerosis [31,32]
- Attenuation of the development of aneurysm induced by angiotensin II [31]
- Reduction of LDL cholesterol levels (-15%) in the PLASMA trial in patients with cardiovascular disease [33]

Phase 2 trials

- Reduction of LDL cholesterol levels (-15%) in the PLASMA trial in patients with cardiovascular disease [33]

Phase 3 trials

- Reduction of Lp-PLA₂-IIA levels and effective in reducing LDL cholesterol levels and C-reactive protein (CRP) in patients with stable and acute cardiovascular disease [23,28]
- VISTA-16 trial in patients with acute coronary syndrome showed an increased incidence of myocardial infarction and of the total of cardiovascular events, mortality, heart attack and stroke [36]

Table 2: Efficacy results of darapladib and varespladib in experimental models and clinical trials

Varespladib Pharmacodynamic properties

- Inhibition of sPLA₂-IIA; IC₅₀=6.2 nM[30]; At similar concentrations it also inhibits sPLA₂-V e X [29]
- 84% inhibition of sPLA₂-IIA at a dose of 500 mg in clinical trials [29]

Experimental models

- Reduction of atherosclerotic plaque area and total cholesterol levels in mouse models of atherosclerosis [31,32]
- Attenuation of the development of aneurysm induced by angiotensin II [31]

Phase 2 trials

- Reduction of LDL cholesterol levels (-15%) in the PLASMA trial in patients with cardiovascular disease [33]

Phase 3 trials

- Reduction of LDL cholesterol levels (-15%) in the PLASMA trial in patients with cardiovascular disease [33]
- 90% reduction in the sPLA-IIa levels and effective in reducing LDL cholesterol levels and C-reactive protein (CRP) in patients with stable and acute cardiovascular disease [23,28]

PLA₂ as Biomarker of Cardiovascular Risk

Epidemiological studies conducted during the past decade have documented that plasma levels of both sPLA₂ and Lp-PLA₂ are correlated with the incidence of cardiovascular disease.

sPLA₂

Two analyses of the epidemiological study EPIC-Norfolk Prospective Population Study showed a significant association between the activity and the mass of sPLA₂-IIA and the onset of the first coronary event. This data was then extended to patients with acute coronary syndrome by demonstrating that a high activity of sPLA₂-
IIA was a predictor factor for further events [19-22]. It should be also noted that a recent meta-analysis, conducted in both the general population and in patients with acute coronary syndrome, showed how the genetic polymorphism in PL2G2A (rs11573156), associated with reduced mass and activity of sPLA2-IIA, was not associated to major cardiovascular events [23], questioning the potential role of this enzyme in acute coronary syndrome.

**PLA2 Associated Lipoprotein**

Regarding the Lp-PLA2, the first evidence of the link between the enzymatic mass and coronary heart disease derived from the population of the case-control study WOSCOPS [24]. Other studies have confirmed this link in a wide spectrum of populations. The results of the PEACE trial also demonstrated that high levels of Lp-PLA2 are indicators of cardiovascular risk in patients with coronary heart disease, independently of traditional risk factors and hs-CRP [25]. Finally, in patients with stroke, the determination of Lp-PLA2 seems to improve the risk stratification [26]. These studies were analyzed in a recent meta-analysis of 32 prospective studies involving 79,036 patients that showed the correlation between activity of Lp-PLA2 and cardiovascular risk [27]. Epidemiological and genetic studies thus represent the most significant evidence of the pro-atherogenic role of Lp-PLA2 and sPLA2-IIA.

**Pharmacological Inhibitors of Lp-LA2 and sPLA2**

The aforementioned experimental, epidemiological, and genetic evidences have provided the rationale for the development of varespladib (Anthera) and darapladib (GSK), sPLA2, and Lp-PLA2 inhibitors, respectively, in the treatment of cardiovascular diseases [28] (Table 2).

The sPLA2 inhibitor varespladib

Varespladib inhibits human sPLA2-IIA, V and X in a powerful and reversible manner [29] (IC50 = 6.2 nM for the IIA) with a selectivity of about 40 times higher than isoform IB [30]. Varespladib was effective in reducing atherosclerosis in murine models of atherosclerosis [31,32], and controlled the development of atheroma induced by angiotensin II [31]. However, in one of these two studies, varespladib reduced levels of total cholesterol, a possible cause of the observed effects. It is interesting to note that, even in the Phase II PLASMA trial (Phospholipase Levels And Serological Markers of Atherosclerosis), varespladib showed a reduction of cholesterol levels in patients with cardiovascular disease [33]. In guinea pigs, an experimental model characterized by the expression of sPLA2-IIA and other isoforms, varespladib treatment did not change the plasma cholesterol levels but reduced the accumulation of lipids in the aortic arch [32]. The observation that varespladib elicits an antiatherosclerotic effect in mice, that do not express the sPLA2-IIA, suggests that the isoforms sPLA2-V,sPLA2-X, or both contribute to the development of atherosclerosis and are inhibited by this drug.

The Lp-PLA2 inhibitor Darapladib

Darapladib is a potent reversible inhibitor of Lp-PLA2 (IC50 = 0.25 nM) [28] which causes a significant reduction in the activity of Lp-PLA2 in atherosclerotic lesions of diabetic/hypercholesterolemic experimental models. Treatment with darapladib reduced the content of lysophosphatidylcholine and attenuated the development of the atherosclerotic plaque [34]. Gene expression analysis has shown an anti-inflammatory effect of darapladib associated with a reduction of the necrotic area [34]. Darapladib has finally demonstrated the ability to reduce the activity of Lp-PLA2 in human carotid plaques [35].

From these clinical and experimental evidences, it is possible to envision some differences between the two pharmacological approaches. The epidemiological evidences of the role of sPLA2 on cardiovascular disease are certainly fewer than those provided for the Lp-PLA2. Particularly, the validity of a therapeutic approach that inhibits the sPLA2-IIA for preventing cardiovascular events has been questioned by the results of a mendelian randomization study [23]. It is also relevant to consider that there are different isoforms of sPLA2, enzymes and only one for the Lp-PLA2. Thus, it is likely that a pharmacological agent would not be able to inhibit all the different sPLA2 isoforms, leading to a possible activation of compensatory sPLA2 activity that could overcome the pharmacological effect. For these reasons, darapladib could be considered a better pharmacological therapy than varespladib. Nevertheless, the advantage of varespladib in comparison to darapladib is potentially due to its effect on plasma cholesterol levels that could contribute to the eventual antatherosclerotic properties.

**Clinical Trials Conducted with Varespladib and Darapladib**

**VISTA-16 trial**

The phase 3 trial VISTA-16 has seen the use of the inhibitor of sPLA2 varespladib in the treatment of patients with acute coronary syndrome [36]. Varespladib not only suppressed the levels of sPLA2-IIA (-90%) but positively affected the lipid-inflammatory status with a significant reduction in the levels of LDL cholesterol and CRP, in patients with both stable and acute cardiovascular (Table 2) [23,28]. Patients enrolled in the VISTA-16 study, treated with 20mg atorvastatin, were randomized within 96 hours after the coronary event to varespladib (500 mg daily) or placebo and stratified according to the cholesterol-lowering therapy and the type of event (STEMI, non-STEMI, unstable angina). The follow-up was 6 months with the visits at 1, 2, 4, 8 and 16 weeks. The results of the study showed an unfavorable effect of varespladib on cardiovascular events, despite the lower LDL cholesterol levels and CRP compared to placebo. Treatment with varespladib caused an increased incidence of myocardial infarction and events of cardiovascular mortality, heart attack and stroke. These results suggest that, inhibition of sPLA2 in the short term with varespladib is harmful in patients with acute coronary syndrome. One possible explanation of the adverse effects observed could be attributed to the fact that varespladib interferes with the pro-atherosclerotic effects of sPLA2-IIA and V but also with anti-atherosclerotic action of isoform X. Although the precise mechanism behind the increase in the incidence of myocardial infarction has not been elucidated, it is possible that varespladib has induced a pro-thrombotic state, although there was no increase in the post-stent thrombosis events. However, it is important to mention that, other drugs modulating the prostaglandin metabolites have shown a detrimental effect on the incidence of myocardial infarction [37]. On the basis of these observations, it is still possible that a selective inhibition of the pro-atherogenic sPLA2 isoforms can exert a favorable action on atherosclerosis.

**Stability trial**

The phase 2 trial IBIS-2, conducted to study the effect of darapladib on plaque stability, showed an arrest in the expansion of the volume of the necrotic core of human carotid plaques assessed by intravascular ultrasound [38]. These findings have led to the hypothesis that darapladib could reduce the risk of cardiovascular events by influencing the composition and the stability of the atherosclerotic plaque. In the phase 3 trial STABILITY (Stabilization of Atherosclerotic Plaque by Darapladib Initiation of Therapy) it has been evaluated the clinical...
efficacy and safety of darapladib in patients with chronic cardiovascular diseases [39]. The study involved the randomization of 15,828 patients to darapladib 160 mg daily or placebo for a period of 3.7 years. All patients were treated according the current guidelines with antidiabetic and statins unless contraindicated or with intolerable side effects. The patients were affected by chronic heart disease, or underwent to a previous myocardial infarction or coronary angioplasty (PCI) or coronary-artery bypass graft (CABG), or had multivascular coronary disease. Darapladib did not significantly reduce the incidence of the primary endpoint, a composite of cardiovascular death, myocardial infarction or stroke. However, there was a significant reduction of the total (a composite of death from coronary heart disease, myocardial infarction, hospitalization for unstable angina, or any coronary revascularization procedure) or major (a composite of death from coronary heart disease, myocardial infarction, or urgent coronary revascularization for myocardial ischemia) coronary events that could be indicative of a possible efficacy of darapladib.

The negative results of this study should be interpreted considering that the patients enrolled were receiving statins, drugs effective in minimizing cardiovascular risk. Indeed, before the randomization, more than a third of the patients had LDL cholesterol levels below 70 mg per deciliter (1.81 mmol/L) and 75% underwent to coronary revascularization. Thus, the conventional therapy has certainly reduced the number of events in the two groups of patients, and therefore those that were potentially modifiable by administration of darapladib.

A second consideration is related to the fact that 96% of patients have received, for the entire duration of the study, statin treatment that can reduce the Lp-PLA₂ levels by 35% [1,2,9]. The effect of statins on Lp-PLA₂ is directly related to their ability to reduce the levels of apoB lipoproteins that carry about 70% of Lp-PLA₂ [29]. Thus, the combination of statins and darapladib could determine an additive cardio protective action through the reduction of Lp-PLA₂ activity. However, the negative results of the study could be explained by at least two reasons: 1) a strong inhibition of Lp-PLA₂ achieved by the combination of statin and darapladib, does not protect from the cardiovascular death, myocardial infarction, or stroke and thus, the pharmacological intervention on Lp-PLA₂ is not an effective therapy for cardiovascular prevention; 2) the effect of darapladib could be masked by the antiatherosclerotic properties of statins or by their anti-inflammatory effects. This second scenario it has been considered relevant also for the development of other antiatherosclerotic drugs that, for ethical reasons, has been tested in clinical trials in patients already under standard of care treatment.

As previously noted, treatment with darapladib resulted in an increase in the incidence of diabetes, along the onset of unpleasant odor of the skin, feces and urine, probably caused by the sulfhydryl group of the molecule of darapladib.

In conclusion, the STABILITY trial has evaluated the effectiveness of a new mechanism for reducing the vulnerability of plaque through the inhibition of Lp-PLA₂, with darapladib in patients with chronic cardiovascular disease previously treated, with conventional therapies. Darapladib did not significantly reduced the primary endpoint of cardiovascular death, myocardial infarction or stroke [39].

**SOLID-TIMI 52 trial**

The therapeutic efficacy of darapladib has also been evaluated in patients with acute coronary syndrome in the phase 3 trial SOLID-TIMI 52 [40]. The study included 13,026 patients, within 30 days of hospitalization for acute coronary syndrome, treated with placebo or darapladib (160 mg daily, 1:1 randomization). Patients were then monitored for an average period of 2.5 years. Similarly to what observed in the STABILITY trial, treatment with darapladib did not alter the primary endpoint of the study of major coronary events (cardiovascular death, myocardial infarction and revascularization). In this case, it has not been observed any favorable effects on secondary endpoints of cardiovascular death, myocardial infarction, stroke, and total mortality. These results were consistent across subgroups including those stratified by baseline LDL cholesterol and the Lp-PLA₂ activity. Although this study did not demonstrate the efficacy of darapladib, there are several limitations to consider. The study was conducted as a fixed dose of 160 mg darapladib capable to inhibit only 66% the Lp-PLA₂ activity, without considering doses more intensive. Furthermore, most of cardiovascular events following the acute coronary syndrome were thrombotic and thus not affected by the treatment with darapladib. In conclusion, these results do not justify the use of Lp-PLA₂ inhibitor darapladib in stable and acute coronary syndrome patients.

**Conclusions**

The negative results of the trials VISTA-16, STABILITY and SOLID-TIMI 52, certainly does not put into question the key role of inflammation in atherogenesis, but emphasize the complexity of the pathogenesis of the atherosclerotic process and the need to further investigate the mechanisms underlying the atherogenesis. The interest in this field is also documented by the fact that there are several anti-inflammatory drugs currently in clinical development. Novartis has just approved the inhibitor of interleukin 1 (IL-1), canakinumab in a phase 3 study on atherosclerosis and the antisense oligonucleotide ISIS-APOARx directed against the lipoprotein (a) would soon start phase 2. Finally, GSK has initiated a phase 3 trial with the p38 MAPK inhibitor (mitogen-activated protein kinase) inhibitor, losmapimod, for a short-term treatment in patients with acute coronary syndrome. Finally, the US National Heart, Lung and Blood Institute have just sponsored a phase 3 trial to evaluate the efficacy of low-dose methotrexate on cardiovascular inflammation. These efforts will help to clarify the efficacy of therapeutic agents with vascular anti-inflammatory activity on the control of atherosclerosis and cardiovascular diseases.

**References**

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