Submit a Manuscript: http://www.wjgnet.com/esps/ Help Desk: http://www.wjgnet.com/esps/helpdesk.aspx DOI: 10.4330/wjc.v9.i1.1 World J Cardiol 2017 January 26; 9(1): 1-13 ISSN 1949-8462 (online) © 2017 Baishideng Publishing Group Inc. All rights reserved.

REVIEW

Endothelial progenitor cells: Exploring the pleiotropic effects of statins

Kully Sandhu, Mamas Mamas, Robert Butler

Kully Sandhu, Mamas Mamas, Robert Butler, Department of Cardiology, Royal Stoke University Hospital, University Hospital of North Midlands, Stafford ST4 6QG, United Kingdom

Author contributions: Sandhu K wrote the paper; Mamas M and Butler R both supervised and reviewed the article.

Conflict-of-interest statement: Authors declare no conflict of interests for this article.

Open-Access: This article is an open-access article which was selected by an in-house editor and fully peer-reviewed by external reviewers. It is distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

Manuscript source: Invited manuscript

Correspondence to: Dr. Kully Sandhu, BSc, DM, MRCP, Cardiology Research Registrar, Department of Cardiology, Royal Stoke University Hospital, University Hospital of North

Midlands, Stoke on Trent, Stafford ST4 6QG, United Kingdom. ksandhu@hotmail.com

Telephone: +44-1782-675953 Fax: +44-1782-672780

Received: June 21, 2016

Peer-review started: June 24, 2016 First decision: August 11, 2016 Revised: September 9, 2016 Accepted: November 1, 2016 Article in press: November 2, 2016 Published online: January 26, 2017

Abstract

Statins have become a cornerstone of risk modification for ischaemic heart disease patients. A number of studies have shown that they are effective and safe.

However studies have observed an early benefit in terms of a reduction in recurrent infarct and or death after a myocardial infarction, prior to any significant change in lipid profile. Therefore, pleiotropic mechanisms, other than lowering lipid profile alone, must account for this effect. One such proposed pleiotropic mechanism is the ability of statins to augment both number and function of endothelial progenitor cells. The ability to augment repair and maintenance of a functioning endothelium may have profound beneficial effect on vascular repair and potentially a positive impact on clinical outcomes in patients with cardiovascular disease. The following literature review will discuss issues surrounding endothelial progenitor cell (EPC) identification, role in vascular repair, factors affecting EPC numbers, the role of statins in current medical practice and their effects on EPC number.

Key words: Statins; Endothelial progenitor cells; Pleiotropic effects; Ischaemic heart disease; Pleiotropic mechanisms

© **The Author(s) 2017.** Published by Baishideng Publishing Group Inc. All rights reserved.

Core tip: Statin therapy is a cornerstone of current management in coronary artery disease. Conventional thinking of stain therapy is for reduction of low-density lipoproteins. However a number of studies have observed an early benefit prior to any significant change in lipid profile. Therefore alternative pleiotropic mechanisms to account for this have been proposed. One such proposed mechanism is the ability of statins to augment both number and function of endothelial progenitor cells (EPCs). The following literature review discusses issues surrounding EPC identification, role in vascular repair, the role of statins in current medical practice and their effects on EPCs.

Sandhu K, Mamas M, Butler R. Endothelial progenitor cells: Exploring the pleiotropic effects of statins. *World J Cardiol*

1



January 26, 2017 | Volume 9 | Issue 1 |

2017; 9(1): 1-13 Available from: URL: http://www.wjgnet.com/1949-8462/full/v9/i1/1.htm DOI: http://dx.doi.org/10.4330/wic.v9.i1.1

INTRODUCTION

The maintenance of endothelial integrity is essential for the preservation of a healthy vasculature^[1]. This integrity results from a balance between on-going endothelial damage and the rate of vascular repair. Disruption of endothelial integrity or impairment of endothelial repair mechanisms is a central step in both the initiation and progression of atherosclerosis^[2]. Endothelial repair is dependent on undifferentiated cells migrating to sites of vascular injury [3-5] then differentiating into mature endothelial cells^[6-13]. These undifferentiated cells are called endothelial progenitor cells (EPCs) have a central role in vascular repair by virtue of their ability to proliferate, migrate to site of vascular injury and then differentiate into mature vascular endothelium^[13,14]. EPCs perpetuate this cycle by secreting pro-angiogenic cytokines[15].

Statins form the corner stone of treatment of coronary artery disease. The safety and efficacy of statins in reducing cardiac events by decreasing serum LDL-cholesterol has been well described^[16-18]. Recently however studies have shown the early beneficial effect of statins occurs before any significant change in lipid profile. This led to the hypothesis that cardiovascular benefits of statins may occur *via* alternative mechanisms other than reduction of LDL-cholesterol alone^[19,20]. One such proposed mechanism is the ability of statins to augment both number and function of EPCs.

The following literature review discusses issues surrounding EPC identification, role in vascular repair, factors affecting EPC numbers, the role of statins in current medical practice and their effects on EPC number.

RESEARCH AND LITERATURE

We performed a review of various studies within the literature available on endothelial progenitor cell and statins. The authors searched various databases (EMBASE, OVID, PubMed) using the keywords: "Endothelial progenitor cells", "statin", "pleiotropic effects". We studied the various publications that we obtained from the search results. Full text manuscripts were obtained. We only included papers in the English language.

EPCS

Cellular identification and staging of differentiation has been made possible by specific surface receptors called epitopes that allow immunophenotyping. This

2

process allows identification of subset of cellular surface molecule termed cluster of differentiation (CD). Cellular subtypes may be defined by the presence or absence of a particular CD molecule. Therefore "CD" may be "+ "or "-" denoting either presence or absence of a particular CD, and is used to describe stem cells rather than fully differentiated cell types. Certain cell types may have variable CD marker expression during maturation for example, and therefore classed as bright (high), mid (mid) or dim (low) denoting intensity of expression^[21,22].

Vascular repair had previously been thought to be due to migration and proliferation of fully differentiated endothelial cells, in a process called angiogenesis^[23]. Asahara et al^[24] identified putative cells with cell surface marker CD34⁺, alternately named kinase insert domain receptor (KDR/VEGFR) markers capable of differentiating into endothelial cells both in vitro and in vivo[24-26]. Subsequent studies recognised that in fact undifferentiated cells subsequently termed EPCs migrated to sites of neovascularization and then differentiate into endothelial cells^[24,26] in a process called vasculogenesis^[27]. EPCs are derived from pluripotential stem cells within the bone marrow. These then evolve into mature endothelial cells[24] accounting for only 0.001%-0.0001% of peripheral blood cells in an unstressed state^[28]. Circulating EPCs may be isolated from bone marrow or the circulation as mononuclear cells^[24,29,30], expressing a variety of endothelial surface markers^[31]. However, there currently remains a lack of consensus on phenotypic and functional definition of endothelial precursor cells[32,33].

EPCs are a diverse group of cells of different lineages that have angiogenic potential, but are not always necessarily able to differentiate into functional endothelial cells as would be suggested from their name^[32]. EPCs are derived from CD34⁺ hematopoietic progenitor cells^[6,24,29,31], with the subset of EPCs characterized by co-expression of endothelial marker proteins^[6,29,31]. Studies have identified 3 markers associated with early functional EPCs including CD133, CD34, and the vascular endothelial growth factor receptor-2 (VEGFR-2) also known as kinase insert domain receptor (KDR, Flk-1 or CD309)^[7,31]. Therefore EPCs express markers of both hematopoietic stem cells (CD34 and CD133) and endothelial cells (CD146, vWF, and VEGFR2)[26,28,29,31,34-37]. The presence of certain cell surface markers are depend on the stage of maturations of the EPC. For example the cell surface marker CD133, a 120-kDa trans-membrane polypeptide, is expressed on bone marrow derived hematopoietic stem and progenitor cells in peripheral blood^[37]. The expression of CD133 on EPCs declines during maturation within the peripheral circulation. Currently there remains some uncertainty as to when EPCs lose the CD133 surface marker, whether during transmigration from bone marrow to circulation or later whilst in the peripheral blood system^[38]. Nevertheless the loss of CD133 represents the transformation into more mature EPCs that are endothelial-like cells^[37].

Table 1 Table to show cell markers during development of endothelial progenitor cell

	Endothelial progenitor cells		
	Bone marrow	Circulation	
		Early EPCs	Mature EPCs
CD133 ⁺	+	+/-	-
CD34⁺	+	+	+
VEGFR2 ⁺	+	++	+++
CD31 ⁺	-	+	+
VE-cadherin	-	+	+
vWF	-	+	+

EPC: Endothelial progenitor cell; VEGFR: Vascular endothelial growth factor receptor.

Whereas the expression of CD34 a cell surface marker found on immature pluripotential stem cells^[31] that acts as an adhesion molecule, although the precise function remains unclear, gradually increases as the CD133 decreases as the EPC matures^[37]. During the course of maturity EPCs begin to have increased expression of other markers specific to endothelial cells such as VEGFR-2, VE-cadherin, and von Willebrand factor (vWF)^[37].

The expression of CD34⁺, CD133⁺, and/or VEGF2⁺ has been used as identifying markers of EPCs in a number of studies^[28,32,39-41]. Whereas as other studies advocate the use of CD133⁺ either alone or in combination with CD34⁺/VEGFR-2⁺ cells for identification of EPCs^[31,42]. In contrast, other studies have suggested that CD133⁺ are haematopoietic cell lines and have not been identified in and therefore unable to form endothelial phenotypic EPCs^[7,40,43]. Ingram *et al*^[28] proposed that CD45⁻ cells incorporated "true" circulating EPCs and verified by other studies^[28,39,40,43]. Interestingly, CD34⁺/VEGFR2⁺ and diminished (dim) CD45 (CD45^{dim}) cells have been found to have greater correlation with coronary heart disease severity and response to statin therapy^[44,45].

In summary, the maturity of the EPC is marked by the gradual loss of CD133, gradual increased expression of CD34⁺ and the appearance of CD31, VE-cadherin and vWF cell surface markers (Table 1).

EPCS AND CORONARY ARTERY DISEASE

Endothelial integrity is essential for healthy vasculature, and can be thought of as a balance between continued endothelial damage and the capacity to repair by a pool of EPCs^[9,46]. It is now generally accepted that cardiovascular risk correlates with EPC numbers. Highlighting the integral relationship between endothelium and atherosclerosis^[47-51], disruption of endothelial integrity by endothelial cell injury has been shown to be a stimulus for the development of atherosclerosis^[2], but also as a stimulus for augmentation of EPC number and function^[9,52,53]. Continued endothelial damage^[54] may lead to an eventual reduction of the number of EPCs. Elevated EPC numbers have been shown to be

associated with augmented formation of collaterals in coronary artery disease^[55] and restoration of endothelial vasodilator function^[9]. A reduction in EPC numbers may lead to deficient endothelial repair and progression of atherosclerosis, with further EPC depletion and perpetuation of atherosclerosis^[9,56]. However, it is uncertain whether low numbers of circulating EPCs represents enhanced usage by vascular repair processes, or reduced production by bone marrow.

CD34⁺ VEGFR2⁺ EPCs cells have been shown to be reduced in patients with atherosclerotic coronary and peripheral disease^[57]. Vasa *et al*, found not only reduced numbers, but also impaired function of EPCs in patients with coronary artery disease. Elevated numbers of EPC have been associated with freedom from myocardial infarction, hospitalization, revascularization and cardiovascular death in patients with coronary artery disease^[56,58]. Furthermore the predictive value of EPC count has been shown to be independent of traditional cardiovascular risk factors^[9,46,59]. In fact, the extent of the reduction in EPC numbers has been associated not only with coronary artery disease burden^[60], but also the presence of symptoms^[61,62].

Finally, elevated numbers of circulating CD34⁺/CD133⁺/VEGFR2 EPCs have been observed after an acute myocardial infarction^[42,63]. This may be regarded as a consequence of cardiac ischaemia together with raised inflammatory and haematopoietic cytokines stimulating EPC mobilisation from the bone marrow^[64-66]. A similar response is seen following coronary angioplasty^[67], and interestingly, the combination of an acute coronary syndrome (ACS) treated by angioplasty provoked an enhanced EPC response^[68]. Therefore, EPC may have a central role not only in repairing coronary vessels after plaque rupture, but also after any coronary intervention.

STATIN THERAPY

Statins act by competitively inhibiting 3 hydroxy-3-methylglutaryl Coenzyme A (HMG CoA) reductase, the rate limiting step in the mevalonate pathway producing isoprenoids including cholesterol. The competitive inhibition of HMG CoA reductase induces the expression of LDL receptors within the liver, thereby increasing the catabolism of plasma LDL, with a consequent decrease in LDL-cholesterol levels^[69].

The safety and efficacy of statins in reducing cardiac events by decreasing serum LDL-cholesterol has been well described^[16-18]. Statin therapy has been shown to reduce death and cardiovascular events in primary prevention of atherosclerosis^[70], stable coronary artery disease^[16,71-73], ACS^[74,75] and secondary prevention^[72]. Statins also appear to reduce development of atherosclerotic lesions^[76,77] and decrease plaque burden^[13,78].

The beneficial effect of intensive statin therapy was studied in a prospective meta-analysis of 90056 patients from 14 randomised trials and found greater

cholesterol reduction was associated with better patient outcomes^[19]. The study found that the 5-year incidence of major adverse cardiac events, coronary revascularization and stroke was reduced by one fifth for every millimoles per liter reduction in LDL cholesterol, which was irrespective of the initial lipid profile^[19].

Another meta-analysis found aggressive statin therapy was associated with reduced peri-procedural myocardial infarction and a 44% risk reduction in major adverse cardiovascular events at 30-d irrespective of clinical presentation^[79]. Moreover, the ARMYDA-RECAPTURE study^[80] found reloading of the high dose statin, atorvastatin 80 mg in 383 NSTEMI and stable angina patients on chronic therapy prior to percutaneous coronary intervention (PCI) had a 50% reduction in 30-d major adverse cardiac events in both group with a greater reduction in NSTEMI group^[80].

These studies led to the universal adoption of statin therapy in patients with coronary artery disease irrespective of presentation from stable angina to ACSs^[81,82].

THE EFFECT OF STATIN THERAPY ON EPCS

Several studies have shown the early beneficial effect of statins occurs before any significant change in lipid profile. This led to the hypothesis that cardiovascular benefits of statins may occur via alternative mechanisms other than reduction of LDL-cholesterol alone^[19,20]. These potential beneficial effect(s) may represent a potential therapeutic target for ischemic heart disease patients, and therefore is of great interest. There have been a number of mechanisms proposed to account for pleiotropic effects of statin therapy. These include reduction in vascular inflammation [83], reduction of platelet aggregability and thrombus deposition[77,84-86], enhancement of fibrinolysis^[87] and increased endothelium derived NO production[88-90]. However the mechanism that has evoked the most interest is the impact of statins on EPCs^[70].

Statin therapy has been associated with greater numbers of circulating EPCs by enhancing mobilization, differentiation, increasing longevity, enhance homing to sites of vascular injury with augmentation of reendothelisation by enhancing expression of adhesion molecules on EPC cell surface membrane^[3,70,91-94].

However, the duration of the effect on EPC number by statin therapy continues to remain contentious.

In one study, atorvastatin therapy was shown to significantly increase circulating EPC as soon as 1 wk with plateauing after 3-4 wk with a 3-fold increase of EPCs from baseline in a stable angina population was also observed^[70]. Whereas Deschaseaux $et\ al^{[95]}$ investigated whether EPCs could be firstly detected and secondly characterized in patients receiving long-term statin therapy defined as 4 wk. The group found a significantly greater number of CD34 $^+$, CD34 $^+$ /CD144 $^+$

circulating EPC in patients receiving statin therapy compared to statin naïve patients. Interestingly two types of EPCs were detected, early and late EPCs. The early EPCs were found to form elongated cells whereas the late EPC population gave rise to cobblestone-like colonies with strong proliferation capacities seen invitro cell culture. The numbers of early EPCs were significantly higher in patients not receiving statin therapy whereas late EPCs were significantly higher in patients receiving statin therapy. The study also observed that long term statin therapy specifically maintained late EPCs in circulation with a CD34⁺/ CD144⁺ phenotype. Rodent studies have found rosuvastatin resulted in a greater than 3 fold increase in EPC numbers when compared with placebo as long as 10 wk after myocardial infarction^[56]. Long-term atorvastatin 10 mg for 12 mo markedly increased EPC number with an associated decrease in oxidative DNA damage^[35]. However to the contrary, Hristov et al^[96] found reduced numbers of circulating EPCs in CHD patients on longterm statin treatment.

Statins appear to have a dose dependant effect on EPC count. A double blinded randomised pilot study found greater number of circulating CD34⁺ VEGFR-2⁺ EPCs after 12 wk of therapy with pravastatin 20 mg when compared to atorvastatin 10 mg^[97]. Similarly, in ACS patients' intensive statin therapy with atorvastatin 80 mg after primary or rescue PCI was associated with greater EPC count at 4-mo follow up as compared to 20 mg atorvastatin. The authors found no beneficial effect in an improvement of LV function^[98]. Furthermore statin reloading in patients on moderate statin therapy undergoing percutaneous coronary intervention has been shown to increase EPC count^[99,100] this correlates with the beneficial effect of statin reloading of high dose statin in patients on chronic therapy^[80] mentioned above.

PLEIOTROPIC EFFECTS OF STATIN THERAPY

Several proposed intracellular signaling mechanisms accounting for the pleiotropic effect of statin therapy have been put forward. Figure 1 below summarizes the positive and negative effects on EPC proliferation, mobilisation and longevity but also the effect of statin therapy.

Nitric oxide pathway

The first proposed intracellular signaling mechanisms involves nitric oxide pathway. The endothelium releases nitric oxide (NO), a primary mediator of smooth muscle tone that causes vasodilatation through the activity of endothelial-type nitric oxide synthase (eNOS)^[101-104]. NO has an central role in vascular homeostasis with its bioavailability dependent on expression of endothelial eNOS^[105], presence of eNOS substrate and or cofactors^[106], phosphorylation of eNOS^[107,108] or due to excessive depletion of NO such as seen with presence of

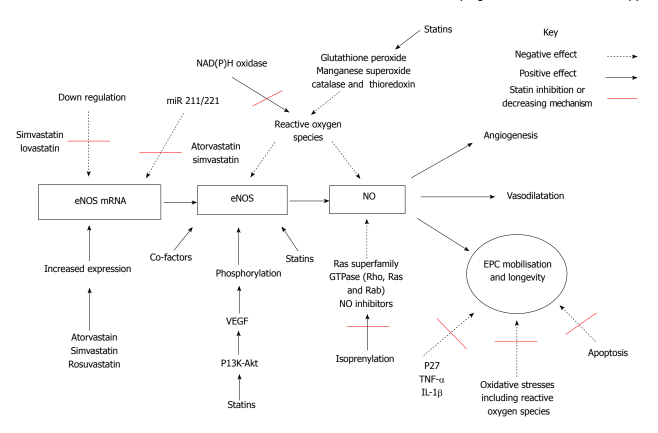


Figure 1 Simplified diagram illustrating the positive and negative effects on endothelial progenitor cell proliferation, mobilisation and longevity together with proposed mechanisms of action of statin therapy. EPC: Endothelial progenitor cell; NO: Nitric oxide; eNOS: Endothelial nitric oxide synthase; VEGF: Vascular endothelial growth factor; mRNA: Messenger ribonucleic acid; TNF: Tumor necrosis factor alpha; IL-1: Interleukin 1; P13k-AKT: Phosphoinositide 3-kinase - protein kinase B pathway; NAD(P)H oxidase: Nicotinamide adenine dinucleotide phosphate-oxidase; miR: Micro non-coding ribonucleic acid.

excessive reactive oxygen species^[109]. However the main functions of NO is as a cellular signaling molecule^[101], an angiogenic factor involved in stimulation, promotion, and stabilization of new blood vessels together with VEGFs, FGFs, Angiopoietins, PDGF, MCP-1, TGF, various integrins, VE-cadherin^[110-113]. Statin therapy has been proposed to both enhance expression and activity of eNOS[114] a prerequisite stage for statin-mediated EPC mobilisation[115]. Statins are known to augment eNOS activity[116-118], increase eNOS expression and restoration of endothelial function[104,119-121]. Statins have also been associated with increased EPC longevity via several pathways including inhibition of p27[122], down regulating TNF- α or IL-1 β expression^[123] and prolonging eNOS expression[122] and finally by increasing eNOS mRNA half-life^[124,125]. Kosmidou *et al*^[126] found simvastatin and rosuvastatin prolonged expression by increasing 3' polyadenylation of eNOS mRNA. Laufs et al^[88,124] firstly noted simvastatin and lovastatin reversed the downregulation of eNOS expression caused by hypoxia and secondly simvastatin reversed down regulation of eNOS expression induced by oxidized LDL[88,124] a recognised cause of atherosclerosis.

miR 221 and miR 222 levels

A second observed pleotropic mechanism of statin therapy has been a decreased level of micro non-coding RNAs called miR 221 and miR 222. These negatively regulate protein expression at post-transcriptional stage[127]. This down regulating effect occurs by targeting 3' untranslated regions resulting in either degradation of target mRNA or impairing translation^[128]. Furthermore miR-221 and miR-222 have been observed to regulate proliferation and differentiation of CD34-positive haematopoietic progenitor cells by reducing expression of c-kit receptor factor impairs haematopoietic progenitor cell proliferation[129]. Increased miR-221 and or miR-221 expression in EPC down regulates EPC differentiation and mobilisation via c-kit and or eNOS pathways in coronary artery disease patients^[127]. Atorvastatin has been shown firstly to decrease miR 221 and miR 222, and secondly increase EPC numbers^[127]. Cerda et al^[130] found both atorvastatin and simvastatin increased NO levels and NOS3 mRNA expression, whereas ezetimibe did not. Atorvastatin, simvastatin and ezetimibe have all been shown to down-regulate the expression of miR-221, whereas miR-222 was reduced only after atorvastatin treatment. The magnitude of the reduction of miR-221 and miR-222 after treatment with statins correlated with an increment in NOS3 mRNA levels[130]. The eNOS and miR221/222 are thought likely to be components of the same pathway[131].

The PI3K/Akt/mTOR pathway

The third proposed pleiotrophic mechanism involves the phosphoinositide 3-kinase (PI3K)/protein kinase B



(Akt)/mammalian target of rapamycin (mTOR) signaling pathway plays. The PI3K/Akt/mTOR pathway plays a central role in multiple cellular processes, including cell proliferation, angiogenesis, metabolism, differentiation and longevity^[132,133]. PI3K generates phosphatidylinositol 3,4,5-triphosphate (PIP3) an important lipid secondary messenger which in turn plays a central role in several signal transduction pathways^[134,135] including activation of the serine/threonine kinases PDKI and Akt. Akt controls protein synthesis and cell growth via the phosphorylation of mTOR^[136]. The PI3K/Akt pathway has been associated with angiogenesis through the regulation of the NO signaling pathway^[137]. The PI3K pathway releases a group of angiogenic factors including VEGF. VEGFR2 has a central role in VEGF-induced angiogenesis $^{\left[138\right] }.$ VEGF is required for the migration of endothelial cells and via PI3K-Akt dependent manner allows formation of capillary like structures^[139]. Studies have shown that NO production may be induced by VEGF and appears to be attenuated by the inhibition of PI3K^[140]. This is thought to occur via phosphorylation of eNOS at the serine 1177 residue by $\mathsf{Akt}^{[107,141]}$, required for the VEGF induced endothelial cell migration^[142]. Factors that stimulate the PI3K/Akt protein kinase pathway, including statins, have been shown also to activate $\text{eNOS}^{[87,\bar{1}41,143]}.$ In turn, the expression of eNOS appears to be fundamental for mobilization of EPC and any impairment in PI3K/Akt/eNOS/NO signaling pathway may result in decreased EPC number^[91,92].

The PI3K/Akt/mTOR intracellular pathway *via* inhibition of the Rho kinase has also been shown to preserves mitochondrial permeability transition pore preventing mitochondrial apoptosis, and therefore death, while conserving cardiomyocyte function^[144,145].

These proposed mechanisms may account for difference in the effect of statin therapy in acute or chronic therapy. Statins given during acute ischaemic stress have been shown to firstly potentiate adenosine receptors^[146,147] eventually leading to downstream regulation of eNOS and therefore increases NO production. Secondly statins augment activation of the reperfusion injury salvage kinase (RISK) pathway^[148]. This results in enhanced activity of the PI3K/Akt/mTOR intracellular signal pathways^[149], leading to preservation of mitochondrial function and cardio-protection. Short-term high dose statin therapy have shown an increase in both EPC mobilisation from bone marrow and augmented function^[92,150-154].

Whereas chronic statin therapy has been linked to a phenomenon termed pre-ischaemic conditioning, protecting the myocardium against ischaemia^[155]. This is believed to be secondary to statin induced NO availability by up regulation of eNOS and stabilisation of eNOS mRNA. Secondly, by increased production of NO and superoxide radicals improves vascular function and reducing vascular inflammation respectively^[88,156]. Statins also inhibit isoprenylation of a number of Ras superfamily GTPase including Rho, Ras and Rab^[157] NO inhibitors resulting in increased NO bioavailability.

Thirdly, by preventing mitochondrial apoptosis and preservation of cardiomyocyte function via the upregulation of the PI3K/Akt/mTOR intracellular signalling pathway by inhibition of Rho kinase^[144,145]. However, the RISK pathway has been shown the down regulated with chronic statin therapy^[158] and has been shown to become reactivated by statin re-loading^[159]. The latter may account for the increase in EPC count in patients on chronic statin therapy reloaded with statin therapy^[80,99,100].

Oxidative stresses

Finally, EPC mobilisation and or function may also be affected by oxidative stress^[153,160]. Oxidative stresses occur secondary to generation of oxygen free radicals or reactive oxygen species (ROS). Oxidative stress has a central role in cardiovascular disease, and a pivotal role in atherosclerosis^[161]. Cellular oxidative stress seen with oxidized low-density lipoprotein (ox-LDL) has a central role in the pathogenesis of atherosclerosis. LDL is oxidised by reactive oxygen species from both circulating cells and cells on vascular walls [162,163]. In essence, LDL oxidation is a result of a chain reaction of free radicals converting polyunsaturated fatty acids into lipid peroxides and as a consequence, formation of active aldehydes^[164]. The biochemical reaction forming ox-LDL have been found to cause senescence of EPCs^[165]. Whereas high density lipoprotein is regarded as atheroprotective due to some part of its antioxidant properties also has a positive effect on EPC number and function^[166]. There are a number of endogenous antioxidants exerting protective effects by scavenging ROS. An indirect way ROS effects EPCs includes ROS reacting with NO forming a potent oxidant $^{[167]}$ with a consequent decrease in NO. Decrease in NO either by excessive oxidation or impaired production reduces EPC mobilisation and/or function^[161,168]. Secondly, direct exposure to oxidative stresses or in disease conditions with high oxidative stress, for example diabetes, is associated with induced EPC apoptosis with significant reduction in EPC numbers [168,169], mobilisation, function[170] and reduced ability to migrate and or integrate into vasculature[161,171].

In an attempt to counteract the effects of oxidative stress EPCs produce superoxide dismutase[172]. Interestingly, cardiovascular risk factors have been found to alter and or reduce the EPC antioxidant ability. Healthy volunteers have found to express higher levels of antioxidative enzyme catalases including glutathione peroxidase and manganese superoxide dismutase when comparing patients with cardiovascular disease^[173,174]. The underlying pathophysiological mechanism currently remains undetermined. The antioxidant pleiotropic effect of statins may include indirect mechanism increasing NO bioavailability accounting for antioxidant properties contributing to an increase in EPC mobilisation and or function^[114,175]. Secondly statin therapy has also been shown to inhibit activation of NAD(P)H oxidase and ROS release^[176] but also activate catalase and thioredoxin

January 26, 2017 | Volume 9 | Issue 1 |

ROS scavenging mechanisms^[176,177]. Finally, statins appear to have a direct effect by significantly reducing peroxide induced apoptosis of EPCs^[169] and decreasing the oxidative damage to DNA in EPCs^[178].

G proteins and G protein-coupled receptors

G protein-coupled receptors (GPCR) are comprised of seven trans-membrane domain proteins and are a super family consisting of a large and diverse number of proteins encoded by approximately 5% of human genes^[179]. There have been a number of classification systems proposed the most recent "GRAFS" (Glutamate, Rhodopsin, Adhesion, Frizzled/Taste2 and Secretin)[180]. In mammals there are five main families $^{[181]}$. GPCRs have an integral role in transfer of extracellular stimuli to within the cell by conformational changes in transmembrane domain structure^[182-185]. They regulate physiological responses to a myriad of endogenous ligands including amines, glycoproteins, peptides and lipids. Therefore, not surprisingly that GPCRs have been implicated in regulation of cellular maintenance, differentiation, proliferation and migration of various stem cells[186-189].

GPCRs modulate activity of intracellular signaling via G proteins. There are currently four known G protein subfamilies each able to potentiate a number of drown stream effectors triggering a number of signaling pathways^[182]. These include activation of Rho associated kinases^[190,191], activation or inhibition of cyclic AMP production^[192] and PI3Ks and therefore modulate the PI3K/Akt pathway^[193,194]. The aforementioned have been implicated in EPC proliferation and function as described above. GPCRs have evoked great interest as a possible target for novel drug therapy [195] as an estimated 50% of all currently prescribed drugs target only a small proportion of GPCRs^[196]. They are also becoming increasingly recognised as having a major role in stem cell signaling^[197]. The role of GPCR in regulation and function of EPCs and the effect of statin therapy remains yet to be elucidated however current evidence suggests that they may have a pivotal role.

CONCLUSION

EPCs have a pivotal role in the maintenance of vascular integrity. However, factors that influence EPC number, migration and function are now becoming recognised and have potentially a significant role in management of ischaemic heart disease patients. Statins once thought to modify cardiovascular risk only by lowering LDL-cholesterol are now being acknowledged as having alternative mechanisms that appear to have beneficial pleiotropic effects. One such mechanism may be mediated by EPCs. A number of studies have shown positive pleiotropic effect of statins on EPCs, both function and number. There appears to be a complex interaction between statins and EPC that is only now becoming recognised. Despite great progress since Asahara's pioneering work, there remain gaps within our

knowledge regarding the pleiotropic effect(s) of statins on EPCs. Further studies are required to elucidate and fully understand any pleiotropic effect and this may guide future beneficial therapeutic interventions.

REFERENCES

- Dong C, Goldschmidt-Clermont PJ. Endothelial progenitor cells: a promising therapeutic alternative for cardiovascular disease. *J Interv Cardiol* 2007; 20: 93-99 [PMID: 17391216 DOI: 10.1111/j.1540-8183.2007.00251.x]
- 2 Ross R. Atherosclerosis--an inflammatory disease. N Engl J Med 1999; 340: 115-126 [PMID: 9887164 DOI: 10.1056/ NEJM199901143400207]
- Walter DH, Rittig K, Bahlmann FH, Kirchmair R, Silver M, Murayama T, Nishimura H, Losordo DW, Asahara T, Isner JM. Statin therapy accelerates reendothelialization: a novel effect involving mobilization and incorporation of bone marrow-derived endothelial progenitor cells. *Circulation* 2002; 105: 3017-3024 [PMID: 12081997 DOI: 10.1161/01.CIR.0000018166.84319.55]
- 4 Griese DP, Ehsan A, Melo LG, Kong D, Zhang L, Mann MJ, Pratt RE, Mulligan RC, Dzau VJ. Isolation and transplantation of autologous circulating endothelial cells into denuded vessels and prosthetic grafts: implications for cell-based vascular therapy. Circulation 2003; 108: 2710-2715 [PMID: 14597586 DOI: 10.1161/01.CIR.0000096490.16596.A6]
- Fujiyama S, Amano K, Uehira K, Yoshida M, Nishiwaki Y, Nozawa Y, Jin D, Takai S, Miyazaki M, Egashira K, Imada T, Iwasaka T, Matsubara H. Bone marrow monocyte lineage cells adhere on injured endothelium in a monocyte chemoattractant protein-1-dependent manner and accelerate reendothelialization as endothelial progenitor cells. Circ Res 2003; 93: 980-989 [PMID: 14525810 DOI: 10.1161/01. RES.0000099245.08637.CE]
- 6 Bhattacharya V, McSweeney PA, Shi Q, Bruno B, Ishida A, Nash R, Storb RF, Sauvage LR, Hammond WP, Wu MH. Enhanced endothelialization and microvessel formation in polyester grafts seeded with CD34(+) bone marrow cells. *Blood* 2000; 95: 581-585 [PMID: 10627466]
- Gehling UM, Ergün S, Schumacher U, Wagener C, Pantel K, Otte M, Schuch G, Schafhausen P, Mende T, Kilic N, Kluge K, Schäfer B, Hossfeld DK, Fiedler W. In vitro differentiation of endothelial cells from AC133-positive progenitor cells. *Blood* 2000; 95: 3106-3112 [PMID: 10807776]
- 8 Hu Y, Davison F, Zhang Z, Xu Q. Endothelial replacement and angiogenesis in arteriosclerotic lesions of allografts are contributed by circulating progenitor cells. *Circulation* 2003; 108: 3122-3127 [PMID: 14656919 DOI: 10.1161/01.CIR.0000105722.96112.67]
- 9 Hill JM, Zalos G, Halcox JP, Schenke WH, Waclawiw MA, Quyyumi AA, Finkel T. Circulating endothelial progenitor cells, vascular function, and cardiovascular risk. N Engl J Med 2003; 348: 593-600 [PMID: 12584367 DOI: 10.1056/NEJMoa022287]
- Takahashi T, Kalka C, Masuda H, Chen D, Silver M, Kearney M, Magner M, Isner JM, Asahara T. Ischemia- and cytokine-induced mobilization of bone marrow-derived endothelial progenitor cells for neovascularization. *Nat Med* 1999; 5: 434-438 [PMID: 10202935 DOI: 10.1038/7434]
- 11 Carmeliet P. Angiogenesis in health and disease. *Nat Med* 2003; 9: 653-660 [PMID: 12778163 DOI: 10.1038/nm0603-653]
- 2 Urbich C, Heeschen C, Aicher A, Dernbach E, Zeiher AM, Dimmeler S. Relevance of monocytic features for neovascularization capacity of circulating endothelial progenitor cells. *Circulation* 2003; 108: 2511-2516 [PMID: 14581410 DOI: 10.1161/01.CIR.00-00096483.29777.50]
- Nissen SE, Nicholls SJ, Sipahi I, Libby P, Raichlen JS, Ballantyne CM, Davignon J, Erbel R, Fruchart JC, Tardif JC, Schoenhagen P, Crowe T, Cain V, Wolski K, Goormastic M, Tuzcu EM. Effect of very high-intensity statin therapy on regression of coronary atherosclerosis: the ASTEROID trial. *JAMA* 2006; 295: 1556-1565 [PMID: 16533939 DOI: 10.1001/jama.295.13.jpc60002]



January 26, 2017 | Volume 9 | Issue 1 |

- 14 Rafii S, Lyden D. Therapeutic stem and progenitor cell transplantation for organ vascularization and regeneration. *Nat Med* 2003; 9: 702-712 [PMID: 12778169 DOI: 10.1038/nm0603-702]
- Mukai N, Akahori T, Komaki M, Li Q, Kanayasu-Toyoda T, Ishii-Watabe A, Kobayashi A, Yamaguchi T, Abe M, Amagasa T, Morita I. A comparison of the tube forming potentials of early and late endothelial progenitor cells. *Exp Cell Res* 2008; 314: 430-440 [PMID: 18083163 DOI: 10.1016/j.yexcr.2007.11.016]
- 16 Randomised trial of cholesterol lowering in 4444 patients with coronary heart disease: the Scandinavian Simvastatin Survival Study (4S). *Lancet* 1994; 344: 1383-1389 [PMID: 7968073]
- Shepherd J, Cobbe SM, Ford I, Isles CG, Lorimer AR, MacFarlane PW, McKillop JH, Packard CJ. Prevention of coronary heart disease with pravastatin in men with hypercholesterolemia. West of Scotland Coronary Prevention Study Group. N Engl J Med 1995; 333: 1301-1307 [PMID: 7566020 DOI: 10.1056/NEJM199511163332001]
- Pedersen TR, Berg K, Cook TJ, Faergeman O, Haghfelt T, Kjekshus J, Miettinen T, Musliner TA, Olsson AG, Pyörälä K, Thorgeirsson G, Tobert JA, Wedel H, Wilhelmsen L. Safety and tolerability of cholesterol lowering with simvastatin during 5 years in the Scandinavian Simvastatin Survival Study. Arch Intern Med 1996; 156: 2085-2092 [PMID: 8862101 DOI: 10.1001/archinte.1996.004401700 97011]
- Baigent C, Keech A, Kearney PM, Blackwell L, Buck G, Pollicino C, Kirby A, Sourjina T, Peto R, Collins R, Simes R. Efficacy and safety of cholesterol-lowering treatment: prospective meta-analysis of data from 90,056 participants in 14 randomised trials of statins. Lancet 2005; 366: 1267-1278 [PMID: 16214597 DOI: 10.1016/S0140-6736(05)67394-1]
- 20 Influence of pravastatin and plasma lipids on clinical events in the West of Scotland Coronary Prevention Study (WOSCOPS). Circulation 1998; 97: 1440-1445 [PMID: 9576423 DOI: 10.1161/01. CIR.97.15.1440]
- 21 Chan JK, Ng CS, Hui PK. A simple guide to the terminology and application of leucocyte monoclonal antibodies. *Histopathology* 1988; 12: 461-480 [PMID: 3294157 DOI: 10.1111/j.1365-2559.1988. tb01967.x]
- 22 Ho IC, Tai TS, Pai SY. GATA3 and the T-cell lineage: essential functions before and after T-helper-2-cell differentiation. *Nat Rev Immunol* 2009; 9: 125-135 [PMID: 19151747 DOI: 10.1038/nri2476]
- 23 Folkman J. Angiogenesis in cancer, vascular, rheumatoid and other disease. *Nat Med* 1995; 1: 27-31 [PMID: 7584949 DOI: 10.1038/ nm0195-27]
- 24 Asahara T, Murohara T, Sullivan A, Silver M, van der Zee R, Li T, Witzenbichler B, Schatteman G, Isner JM. Isolation of putative progenitor endothelial cells for angiogenesis. *Science* 1997; 275: 964-967 [PMID: 9020076 DOI: 10.1126/science.275.5302.964]
- 25 Stump MM, Jordan GL, Debakey ME, Halpert B. Endothelium grown from circulating blood on isolated intravascular dacron hub. Am J Pathol 1963; 43: 361-367 [PMID: 14057632]
- 26 Asahara T, Masuda H, Takahashi T, Kalka C, Pastore C, Silver M, Kearne M, Magner M, Isner JM. Bone marrow origin of endothelial progenitor cells responsible for postnatal vasculogenesis in physiological and pathological neovascularization. *Circ Res* 1999; 85: 221-228 [PMID: 10436164 DOI: 10.1161/01.RES.85.3.221]
- 27 Risau W. Mechanisms of angiogenesis. *Nature* 1997; 386: 671-674 [PMID: 9109485 DOI: 10.1038/386671a0]
- 28 Ingram DA, Caplice NM, Yoder MC. Unresolved questions, changing definitions, and novel paradigms for defining endothelial progenitor cells. *Blood* 2005; 106: 1525-1531 [PMID: 15905185 DOI: 10.1182/blood-2005-04-1509]
- 29 Shi Q, Rafii S, Wu MH, Wijelath ES, Yu C, Ishida A, Fujita Y, Kothari S, Mohle R, Sauvage LR, Moore MA, Storb RF, Hammond WP. Evidence for circulating bone marrow-derived endothelial cells. *Blood* 1998; 92: 362-367 [PMID: 9657732]
- 30 Lin Y, Weisdorf DJ, Solovey A, Hebbel RP. Origins of circulating endothelial cells and endothelial outgrowth from blood. J Clin Invest 2000; 105: 71-77 [PMID: 10619863 DOI: 10.1172/JCI8071]
- 31 Peichev M, Naiyer AJ, Pereira D, Zhu Z, Lane WJ, Williams M,

- Oz MC, Hicklin DJ, Witte L, Moore MA, Rafii S. Expression of VEGFR-2 and AC133 by circulating human CD34(+) cells identifies a population of functional endothelial precursors. *Blood* 2000; **95**: 952-958 [PMID: 10648408]
- 32 Hirschi KK, Ingram DA, Yoder MC. Assessing identity, phenotype, and fate of endothelial progenitor cells. *Arterioscler Thromb Vasc Biol* 2008; 28: 1584-1595 [PMID: 18669889 DOI: 10.1161/ATVBAHA.107.155960]
- Fadini GP, Losordo D, Dimmeler S. Critical reevaluation of endothelial progenitor cell phenotypes for therapeutic and diagnostic use. *Circ Res* 2012; 110: 624-637 [PMID: 22343557 DOI: 10.1161/ CIRCRESAHA.111.243386]
- 34 Fadini GP, Baesso I, Albiero M, Sartore S, Agostini C, Avogaro A. Technical notes on endothelial progenitor cells: ways to escape from the knowledge plateau. *Atherosclerosis* 2008; 197: 496-503 [PMID: 18249408 DOI: 10.1016/j.atherosclerosis.2007.12.039]
- 35 Hristov M, Erl W, Weber PC. Endothelial progenitor cells: mobilization, differentiation, and homing. *Arterioscler Thromb Vasc Biol* 2003; 23: 1185-1189 [PMID: 12714439 DOI: 10.1161/01. ATV.0000073832.49290.B5]
- 36 Urbich C, Dimmeler S. Endothelial progenitor cells: characterization and role in vascular biology. *Circ Res* 2004; 95: 343-353 [PMID: 15321944 DOI: 10.1161/01.RES.0000137877.89448.78]
- 37 Yin AH, Miraglia S, Zanjani ED, Almeida-Porada G, Ogawa M, Leary AG, Olweus J, Kearney J, Buck DW. AC133, a novel marker for human hematopoietic stem and progenitor cells. *Blood* 1997; 90: 5002-5012 [PMID: 9389720]
- Quirici N, Soligo D, Caneva L, Servida F, Bossolasco P, Deliliers GL. Differentiation and expansion of endothelial cells from human bone marrow CD133(+) cells. *Br J Haematol* 2001; 115: 186-194 [PMID: 11722432 DOI: 10.1046/j.1365-2141.2001.03077.x]
- 39 Bertolini F, Shaked Y, Mancuso P, Kerbel RS. The multifaceted circulating endothelial cell in cancer: towards marker and target identification. *Nat Rev Cancer* 2006; 6: 835-845 [PMID: 17036040 DOI: 10.1038/nrc1971]
- 40 Timmermans F, Van Hauwermeiren F, De Smedt M, Raedt R, Plasschaert F, De Buyzere ML, Gillebert TC, Plum J, Vandekerckhove B. Endothelial outgrowth cells are not derived from CD133+cells or CD45+ hematopoietic precursors. *Arterioscler Thromb Vasc Biol* 2007; 27: 1572-1579 [PMID: 17495235 DOI: 10.1161/ATVBAHA.107.144972]
- 41 Friedrich EB, Walenta K, Scharlau J, Nickenig G, Werner N. CD34-/ CD133+/VEGFR-2+ endothelial progenitor cell subpopulation with potent vasoregenerative capacities. *Circ Res* 2006; 98: e20-e25 [PMID: 16439688 DOI: 10.1161/01.RES.0000205765.28940.93]
- 42 Gill M, Dias S, Hattori K, Rivera ML, Hicklin D, Witte L, Girardi L, Yurt R, Himel H, Rafii S. Vascular trauma induces rapid but transient mobilization of VEGFR2(+)AC133(+) endothelial precursor cells. Circ Res 2001; 88: 167-174 [PMID: 11157668 DOI: 10.1161/01.RES.88.2.167]
- 43 Case J, Mead LE, Bessler WK, Prater D, White HA, Saadatzadeh MR, Bhavsar JR, Yoder MC, Haneline LS, Ingram DA. Human CD34+AC133+VEGFR-2+ cells are not endothelial progenitor cells but distinct, primitive hematopoietic progenitors. *Exp Hematol* 2007; 35: 1109-1118 [PMID: 17588480 DOI: 10.1016/j.exphem.2007.04.002]
- 44 Schmidt-Lucke C, Fichtlscherer S, Aicher A, Tschöpe C, Schultheiss HP, Zeiher AM, Dimmeler S. Quantification of circulating endothelial progenitor cells using the modified ISHAGE protocol. *PLoS One* 2010; 5: e13790 [PMID: 21072182 DOI: 10.1371/journal. pone.0013790]
- 45 Farace F, Gross-Goupil M, Tournay E, Taylor M, Vimond N, Jacques N, Billiot F, Mauguen A, Hill C, Escudier B. Levels of circulating CD 45(dim)CD34(+)VEGFR2(+) progenitor cells correlate with outcome in metastatic renal cell carcinoma patients treated with tyrosine kinase inhibitors. *Br J Cancer* 2011; 104: 1144-1150 [PMID: 21386843 DOI: 10.1038/bjc.2011.72]
- 46 Schmidt-Lucke C, Rössig L, Fichtlscherer S, Vasa M, Britten M, Kämper U, Dimmeler S, Zeiher AM. Reduced number of circulating endothelial progenitor cells predicts future cardiovascular events:



- proof of concept for the clinical importance of endogenous vascular repair. *Circulation* 2005; **111**: 2981-2987 [PMID: 15927972 DOI: 10.1161/CIRCULATIONAHA.104.504340]
- 47 Padfield GJ, Newby DE, Mills NL. Understanding the role of endothelial progenitor cells in percutaneous coronary intervention. *J Am Coll Cardiol* 2010; 55: 1553-1565 [PMID: 20378071 DOI: 10.1016/j.jacc.2009.10.070]
- 48 Schächinger V, Britten MB, Zeiher AM. Prognostic impact of coronary vasodilator dysfunction on adverse long-term outcome of coronary heart disease. *Circulation* 2000; 101: 1899-1906 [PMID: 10779454 DOI: 10.1161/01.CIR.101.16.1899]
- 49 Suwaidi JA, Hamasaki S, Higano ST, Nishimura RA, Holmes DR, Lerman A. Long-term follow-up of patients with mild coronary artery disease and endothelial dysfunction. *Circulation* 2000; 101: 948-954 [PMID: 10704159 DOI: 10.1161/01.CIR.101.9.948]
- 50 Halcox JP, Schenke WH, Zalos G, Mincemoyer R, Prasad A, Waclawiw MA, Nour KR, Quyyumi AA. Prognostic value of coronary vascular endothelial dysfunction. *Circulation* 2002; 106: 653-658 [PMID: 12163423 DOI: 10.1161/01.CIR.0000025404.78001.D8]
- 51 Gokce N, Keaney JF, Hunter LM, Watkins MT, Menzoian JO, Vita JA. Risk stratification for postoperative cardiovascular events via noninvasive assessment of endothelial function: a prospective study. *Circulation* 2002; 105: 1567-1572 [PMID: 11927524 DOI: 10.1161/01.CIR.0000012543.55874.47]
- 52 Rouhl RP, van Oostenbrugge RJ, Damoiseaux J, Tervaert JW, Lodder J. Endothelial progenitor cell research in stroke: a potential shift in pathophysiological and therapeutical concepts. Stroke 2008; 39: 2158-2165 [PMID: 18451350 DOI: 10.1161/STROKEAHA.107.507251]
- 53 Sen S, McDonald SP, Coates PT, Bonder CS. Endothelial progenitor cells: novel biomarker and promising cell therapy for cardiovascular disease. *Clin Sci* (Lond) 2011; 120: 263-283 [PMID: 21143202 DOI: 10.1042/CS20100429]
- 54 Dimmeler S, Vasa-Nicotera M. Aging of progenitor cells: limitation for regenerative capacity? *J Am Coll Cardiol* 2003; 42: 2081-2082 [PMID: 14680730 DOI: 10.1016/j.jacc.2003.09.016]
- 55 Lambiase PD, Edwards RJ, Anthopoulos P, Rahman S, Meng YG, Bucknall CA, Redwood SR, Pearson JD, Marber MS. Circulating humoral factors and endothelial progenitor cells in patients with differing coronary collateral support. *Circulation* 2004; 109: 2986-2992 [PMID: 15184289 DOI: 10.1161/01.CIR.0000130639.97284.EC]
- Vasa M, Fichtlscherer S, Aicher A, Adler K, Urbich C, Martin H, Zeiher AM, Dimmeler S. Number and migratory activity of circulating endothelial progenitor cells inversely correlate with risk factors for coronary artery disease. *Circ Res* 2001; 89: E1-E7 [PMID: 11440984 DOI: 10.1161/hh1301.093953]
- 57 Chironi G, Walch L, Pernollet MG, Gariepy J, Levenson J, Rendu F, Simon A. Decreased number of circulating CD34+KDR+ cells in asymptomatic subjects with preclinical atherosclerosis. *Atherosclerosis* 2007; 191: 115-120 [PMID: 16620831 DOI: 10.1016/j.atherosclerosis. 2006.02.041]
- Werner N, Wassmann S, Ahlers P, Schiegl T, Kosiol S, Link A, Walenta K, Nickenig G. Endothelial progenitor cells correlate with endothelial function in patients with coronary artery disease. *Basic Res Cardiol* 2007; 102: 565-571 [PMID: 17932708 DOI: 10.1007/s00395-007-0680-1]
- Werner N, Kosiol S, Schiegl T, Ahlers P, Walenta K, Link A, Böhm M, Nickenig G. Circulating endothelial progenitor cells and cardiovascular outcomes. *N Engl J Med* 2005; 353: 999-1007 [PMID: 16148285 DOI: 10.1056/NEJMoa043814]
- 60 Kunz GA, Liang G, Cuculi F, Gregg D, Vata KC, Shaw LK, Goldschmidt-Clermont PJ, Dong C, Taylor DA, Peterson ED. Circulating endothelial progenitor cells predict coronary artery disease severity. *Am Heart J* 2006; **152**: 190-195 [PMID: 16824855 DOI: 10.1016/j.ahj.2006.02.001]
- 61 Gaspardone A, Menghini F, Mazzuca V, Skossyreva O, Barbato G, de Fabritiis P. Progenitor cell mobilisation in patients with acute and chronic coronary artery disease. *Heart* 2006; 92: 253-254 [PMID: 16415195 DOI: 10.1136/hrt.2004.058636]
- 62 Güven H, Shepherd RM, Bach RG, Capoccia BJ, Link DC. The

- number of endothelial progenitor cell colonies in the blood is increased in patients with angiographically significant coronary artery disease. *J Am Coll Cardiol* 2006; **48**: 1579-1587 [PMID: 17045891 DOI: 10.1016/j.jacc.2006.04.101]
- 63 Shintani S, Murohara T, Ikeda H, Ueno T, Honma T, Katoh A, Sasaki K, Shimada T, Oike Y, Imaizumi T. Mobilization of endothelial progenitor cells in patients with acute myocardial infarction. *Circulation* 2001; 103: 2776-2779 [PMID: 11401930 DOI: 10.1161/hc2301.092122]
- 64 Wojakowski W, Tendera M, Michałowska A, Majka M, Kucia M, Maślankiewicz K, Wyderka R, Ochała A, Ratajczak MZ. Mobilization of CD34/CXCR4+, CD34/CD117+, c-met+ stem cells, and mononuclear cells expressing early cardiac, muscle, and endothelial markers into peripheral blood in patients with acute myocardial infarction. *Circulation* 2004; 110: 3213-3220 [PMID: 15533859 DOI: 10.1161/01.CIR.0000147609.39780.02]
- Massa M, Rosti V, Ferrario M, Campanelli R, Ramajoli I, Rosso R, De Ferrari GM, Ferlini M, Goffredo L, Bertoletti A, Klersy C, Pecci A, Moratti R, Tavazzi L. Increased circulating hematopoietic and endothelial progenitor cells in the early phase of acute myocardial infarction. *Blood* 2005; 105: 199-206 [PMID: 15345590 DOI: 10.1182/blood-2004-05-1831]
- 66 Leone AM, Rutella S, Bonanno G, Abbate A, Rebuzzi AG, Giovannini S, Lombardi M, Galiuto L, Liuzzo G, Andreotti F, Lanza GA, Contemi AM, Leone G, Crea F. Mobilization of bone marrow-derived stem cells after myocardial infarction and left ventricular function. Eur Heart J 2005; 26: 1196-1204 [PMID: 15734770 DOI: 10.1093/eurheartj/ehi164]
- 67 Banerjee S, Brilakis E, Zhang S, Roesle M, Lindsey J, Philips B, Blewett CG, Terada LS. Endothelial progenitor cell mobilization after percutaneous coronary intervention. *Atherosclerosis* 2006; 189: 70-75 [PMID: 16806234 DOI: 10.1016/j.atherosclerosis.2006. 04.026]
- 68 Laufs U, Liao JK. Targeting Rho in cardiovascular disease. Circ Res 2000; 87: 526-528 [PMID: 11009552 DOI: 10.1161/01. RES.87.7.526]
- 69 Istvan ES. Structural mechanism for statin inhibition of 3-hydroxy-3-methylglutaryl coenzyme A reductase. *Am Heart J* 2002; 144: S27-S32 [PMID: 12486413 DOI: 10.1067/mhj.2002.130300]
- Vasa M, Fichtlscherer S, Adler K, Aicher A, Martin H, Zeiher AM, Dimmeler S. Increase in circulating endothelial progenitor cells by statin therapy in patients with stable coronary artery disease. Circulation 2001; 103: 2885-2890 [PMID: 11413075 DOI: 10.1161/hc2401.092816]
- Prevention of cardiovascular events and death with pravastatin in patients with coronary heart disease and a broad range of initial cholesterol levels. The Long-Term Intervention with Pravastatin in Ischaemic Disease (LIPID) Study Group. N Engl J Med 1998; 339: 1349-1357 [PMID: 9841303 DOI: 10.1056/NEJM199811053391902]
- 72 Sacks FM, Pfeffer MA, Moye LA, Rouleau JL, Rutherford JD, Cole TG, Brown L, Warnica JW, Arnold JM, Wun CC, Davis BR, Braunwald E. The effect of pravastatin on coronary events after myocardial infarction in patients with average cholesterol levels. Cholesterol and Recurrent Events Trial investigators. N Engl J Med 1996; 335: 1001-1009 [PMID: 8801446 DOI: 10.1056/NEJM199610033351401]
- 73 LaRosa JC, Grundy SM, Waters DD, Shear C, Barter P, Fruchart JC, Gotto AM, Greten H, Kastelein JJ, Shepherd J, Wenger NK. Intensive lipid lowering with atorvastatin in patients with stable coronary disease. N Engl J Med 2005; 352: 1425-1435 [PMID: 15755765 DOI: 10.1056/NEJMoa050461]
- Ridker PM, Morrow DA, Rose LM, Rifai N, Cannon CP, Braunwald E. Relative efficacy of atorvastatin 80 mg and pravastatin 40 mg in achieving the dual goals of low-density lipoprotein cholesterol & It; 70 mg/dl and C-reactive protein & It; 2 mg/l: an analysis of the PROVE-IT TIMI-22 trial. *J Am Coll Cardiol* 2005; 45: 1644-1648 [PMID: 15893181 DOI: 10.1016/j.jacc.2005.02.080]
- 75 Schwartz GG, Olsson AG, Ezekowitz MD, Ganz P, Oliver MF, Waters D, Zeiher A, Chaitman BR, Leslie S, Stern T. Effects of atorvastatin on early recurrent ischemic events in acute coronary



January 26, 2017 | Volume 9 | Issue 1

- syndromes: the MIRACL study: a randomized controlled trial. *JAMA* 2001; **285**: 1711-1718 [PMID: 11277825 DOI: 10.1001/jama 285 13 1711]
- 76 Maron DJ, Fazio S, Linton MF. Current perspectives on statins. Circulation 2000; 101: 207-213 [PMID: 10637210 DOI: 10.1161/01. CIR.101.2.207]
- 77 Lacoste L, Lam JY, Hung J, Letchacovski G, Solymoss CB, Waters D. Hyperlipidemia and coronary disease. Correction of the increased thrombogenic potential with cholesterol reduction. *Circulation* 1995; 92: 3172-3177 [PMID: 7586300 DOI: 10.1161/01.CIR.92.11.3172]
- Nissen SE, Tuzcu EM, Schoenhagen P, Brown BG, Ganz P, Vogel RA, Crowe T, Howard G, Cooper CJ, Brodie B, Grines CL, DeMaria AN. Effect of intensive compared with moderate lipid-lowering therapy on progression of coronary atherosclerosis: a randomized controlled trial. *JAMA* 2004; 291: 1071-1080 [PMID: 14996776 DOI: 10.1001/jama.291.9.1071]
- 79 Patti G, Cannon CP, Murphy SA, Mega S, Pasceri V, Briguori C, Colombo A, Yun KH, Jeong MH, Kim JS, Choi D, Bozbas H, Kinoshita M, Fukuda K, Jia XW, Hara H, Cay S, Di Sciascio G. Clinical benefit of statin pretreatment in patients undergoing percutaneous coronary intervention: a collaborative patient-level meta-analysis of 13 randomized studies. *Circulation* 2011; 123: 1622-1632 [PMID: 21464051 DOI: 10.1161/CIRCULATIONAHA.110.002451]
- 80 Di Sciascio G, Patti G, Pasceri V, Gaspardone A, Colonna G, Montinaro A. Efficacy of atorvastatin reload in patients on chronic statin therapy undergoing percutaneous coronary intervention: results of the ARMYDA-RECAPTURE (Atorvastatin for Reduction of Myocardial Damage During Angioplasty) Randomized Trial. J Am Coll Cardiol 2009; 54: 558-565 [PMID: 19643320 DOI: 10.1016/j.jacc.2009.05.028]
- 81 Amsterdam EA, Wenger NK, Brindis RG, Casey DE, Ganiats TG, Holmes DR, Jaffe AS, Jneid H, Kelly RF, Kontos MC, Levine GN, Liebson PR, Mukherjee D, Peterson ED, Sabatine MS, Smalling RW, Zieman SJ. 2014 AHA/ACC Guideline for the Management of Patients with Non-ST-Elevation Acute Coronary Syndromes: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol* 2014; 64: e139-e228 [PMID: 25260718 DOI: 10.1016/j.jacc.2014.09.017]
- 82 O'Gara PT, Kushner FG, Ascheim DD, Casey DE, Chung MK, de Lemos JA, Ettinger SM, Fang JC, Fesmire FM, Franklin BA, Granger CB, Krumholz HM, Linderbaum JA, Morrow DA, Newby LK, Ornato JP, Ou N, Radford MJ, Tamis-Holland JE, Tommaso CL, Tracy CM, Woo YJ, Zhao DX, Anderson JL, Jacobs AK, Halperin JL, Albert NM, Brindis RG, Creager MA, DeMets D, Guyton RA, Hochman JS, Kovacs RJ, Kushner FG, Ohman EM, Stevenson WG, Yancy CW. 2013 ACCF/AHA guideline for the management of ST-elevation myocardial infarction: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation 2013; 127: e362-e425 [PMID: 23247304 DOI: 10.1161/CIR.0b013e3182742cf6]
- 83 Bustos C, Hernández-Presa MA, Ortego M, Tuñón J, Ortega L, Pérez F, Díaz C, Hernández G, Egido J. HMG-CoA reductase inhibition by atorvastatin reduces neointimal inflammation in a rabbit model of atherosclerosis. *J Am Coll Cardiol* 1998; 32: 2057-2064 [PMID: 9857893 DOI: 10.1016/S0735-1097(98)00487-2]
- 84 Laufs U, Gertz K, Huang P, Nickenig G, Böhm M, Dirnagl U, Endres M. Atorvastatin upregulates type III nitric oxide synthase in thrombocytes, decreases platelet activation, and protects from cerebral ischemia in normocholesterolemic mice. *Stroke* 2000; 31: 2442-2449 [PMID: 11022078 DOI: 10.1161/01.STR.31.10.2442]
- 85 Dogra GK, Watts GF, Chan DC, Stanton K. Statin therapy improves brachial artery vasodilator function in patients with Type 1 diabetes and microalbuminuria. *Diabet Med* 2005; 22: 239-242 [PMID: 15717868 DOI: 10.1111/j.1464-5491.2004.01382.x]
- 86 Dangas G, Smith DA, Unger AH, Shao JH, Meraj P, Fier C, Cohen AM, Fallon JT, Badimon JJ, Ambrose JA. Pravastatin: an antithrombotic effect independent of the cholesterol-lowering effect. Thromb Haemost 2000; 83: 688-692 [PMID: 10823263]

- 87 Dimmeler S, Aicher A, Vasa M, Mildner-Rihm C, Adler K, Tiemann M, Rütten H, Fichtlscherer S, Martin H, Zeiher AM. HMG-CoA reductase inhibitors (statins) increase endothelial progenitor cells via the PI 3-kinase/Akt pathway. *J Clin Invest* 2001; 108: 391-397 [PMID: 11489932 DOI: 10.1172/JCI13152]
- Laufs U, La Fata V, Plutzky J, Liao JK. Upregulation of endothelial nitric oxide synthase by HMG CoA reductase inhibitors. *Circulation* 1998; 97: 1129-1135 [PMID: 9537338 DOI: 10.1161/01. CIR 97.12.1129]
- 89 Kureishi Y, Luo Z, Shiojima I, Bialik A, Fulton D, Lefer DJ, Sessa WC, Walsh K. The HMG-CoA reductase inhibitor simvastatin activates the protein kinase Akt and promotes angiogenesis in normocholesterolemic animals. *Nat Med* 2000; 6: 1004-1010 [PMID: 10973320 DOI: 10.1038/79510]
- 90 Laufs U, Liao JK. Post-transcriptional regulation of endothelial nitric oxide synthase mRNA stability by Rho GTPase. *J Biol Chem* 1998; 273: 24266-24271 [PMID: 9727051 DOI: 10.1074/jbc.273.37.24266]
- 91 Aicher A, Heeschen C, Mildner-Rihm C, Urbich C, Ihling C, Technau-Ihling K, Zeiher AM, Dimmeler S. Essential role of endothelial nitric oxide synthase for mobilization of stem and progenitor cells. *Nat Med* 2003; 9: 1370-1376 [PMID: 14556003 DOI: 10.1038/nm948]
- 92 Llevadot J, Murasawa S, Kureishi Y, Uchida S, Masuda H, Kawamoto A, Walsh K, Isner JM, Asahara T. HMG-CoA reductase inhibitor mobilizes bone marrow--derived endothelial progenitor cells. *J Clin Invest* 2001; 108: 399-405 [PMID: 11489933 DOI: 10.1172/JCI13131]
- 93 Bush LR, Shebuski RJ. In vivo models of arterial thrombosis and thrombolysis. FASEB J 1990; 4: 3087-3098 [PMID: 2210155]
- 94 Urbich C, Dernbach E, Zeiher AM, Dimmeler S. Double-edged role of statins in angiogenesis signaling. *Circ Res* 2002; 90: 737-744 [PMID: 11934843 DOI: 10.1161/01.RES.0000014081.30867.F8]
- 95 Deschaseaux F, Selmani Z, Falcoz PE, Mersin N, Meneveau N, Penfornis A, Kleinclauss C, Chocron S, Etievent JP, Tiberghien P, Kantelip JP, Davani S. Two types of circulating endothelial progenitor cells in patients receiving long term therapy by HMG-CoA reductase inhibitors. Eur J Pharmacol 2007; 562: 111-118 [PMID: 17320859 DOI: 10.1016/j.ejphar.2007.01.045]
- 96 Hristov M, Fach C, Becker C, Heussen N, Liehn EA, Blindt R, Hanrath P, Weber C. Reduced numbers of circulating endothelial progenitor cells in patients with coronary artery disease associated with long-term statin treatment. *Atherosclerosis* 2007; 192: 413-420 [PMID: 16837000 DOI: 10.1016/j.atherosclerosis.2006.05.031]
- 97 Lin LY, Huang CC, Chen JS, Wu TC, Leu HB, Huang PH, Chang TT, Lin SJ, Chen JW. Effects of pitavastatin versus atorvastatin on the peripheral endothelial progenitor cells and vascular endothelial growth factor in high-risk patients: a pilot prospective, double-blind, randomized study. *Cardiovasc Diabetol* 2014; 13: 111 [PMID: 25027585 DOI: 10.1186/s12933-014-0111-1]
- 98 Leone AM, Rutella S, Giannico MB, Perfetti M, Zaccone V, Brugaletta S, Garramone B, Niccoli G, Porto I, Liuzzo G, Biasucci LM, Bellesi S, Galiuto L, Leone G, Rebuzzi AG, Crea F. Effect of intensive vs standard statin therapy on endothelial progenitor cells and left ventricular function in patients with acute myocardial infarction: Statins for regeneration after acute myocardial infarction and PCI (STRAP) trial. *Int J Cardiol* 2008; 130: 457-462 [PMID: 18667247 DOI: 10.1016/j.ijcard.2008.05.036]
- Ye H, He F, Fei X, Lou Y, Wang S, Yang R, Hu Y, Chen X. High-dose atorvastatin reloading before percutaneous coronary intervention increased circulating endothelial progenitor cells and reduced inflammatory cytokine expression during the perioperative period. *J Cardiovasc Pharmacol Ther* 2014; 19: 290-295 [PMID: 24346155 DOI: 10.1177/1074248413513500]
- 100 Ricottini EMR, Patti G. Benefit of atorvastatin reload on endothelial progenitor cells in patients on chronic statin treatment undergoing PCI. *J Am Coll Cardiol* 2013; 61: 10 [DOI: 10.1016/S0735-1097(13)61635-6]
- 101 Knowles RG, Moncada S. Nitric oxide synthases in mammals. Biochem J 1994; 298 (Pt 2): 249-258 [PMID: 7510950 DOI:



- 10.1042/bj2980249]
- 102 Moncada S, Higgs A. The L-arginine-nitric oxide pathway. N Engl J Med 1993; 329: 2002-2012 [PMID: 7504210 DOI: 10.1056/ NEJM199312303292706]
- 103 Sessa WC. The nitric oxide synthase family of proteins. *J Vasc Res* 1994; 31: 131-143 [PMID: 7511942 DOI: 10.1159/000159039]
- 104 Moncada S, Palmer RM, Higgs EA. Nitric oxide: physiology, pathophysiology, and pharmacology. *Pharmacol Rev* 1991; 43: 109-142 [PMID: 1852778]
- 105 Wilcox JN, Subramanian RR, Sundell CL, Tracey WR, Pollock JS, Harrison DG, Marsden PA. Expression of multiple isoforms of nitric oxide synthase in normal and atherosclerotic vessels. *Arterioscler Thromb Vasc Biol* 1997; 17: 2479-2488 [PMID: 9409218 DOI: 10.1161/01.ATV.17.11.2479]
- 106 Stuehr DJ, Griffith OW. Mammalian nitric oxide synthases. *Adv Enzymol Relat Areas Mol Biol* 1992; 65: 287-346 [PMID: 1373932 DOI: 10.1002/9780470123119.ch8]
- 107 Dimmeler S, Fleming I, Fisslthaler B, Hermann C, Busse R, Zeiher AM. Activation of nitric oxide synthase in endothelial cells by Akt-dependent phosphorylation. *Nature* 1999; 399: 601-605 [PMID: 10376603 DOI: 10.1038/21224]
- 108 Fleming I, Fisslthaler B, Dimmeler S, Kemp BE, Busse R. Phosphorylation of Thr(495) regulates Ca(2+)/calmodulin-dependent endothelial nitric oxide synthase activity. *Circ Res* 2001; 88: E68-E75 [PMID: 11397791 DOI: 10.1161/hh1101.092677]
- 109 Rubanyi GM, Vanhoutte PM. Superoxide anions and hyperoxia inactivate endothelium-derived relaxing factor. Am J Physiol 1986; 250: H822-H827 [PMID: 3010744]
- 110 Carmeliet P. Mechanisms of angiogenesis and arteriogenesis. *Nat Med* 2000; 6: 389-395 [PMID: 10742145 DOI: 10.1038/74651]
- 111 Carmeliet P, Jain RK. Angiogenesis in cancer and other diseases. *Nature* 2000; 407: 249-257 [PMID: 11001068 DOI: 10.1038/35025220]
- 112 Semenza GL. Vasculogenesis, angiogenesis, and arteriogenesis: mechanisms of blood vessel formation and remodeling. *J Cell Biochem* 2007; 102: 840-847 [PMID: 17891779 DOI: 10.1002/jcb.21523]
- 113 Wahlberg E. Angiogenesis and arteriogenesis in limb ischemia. J Vasc Surg 2003; 38: 198-203 [PMID: 12844115 DOI: 10.1016/ S0741-5214(03)00151-4]
- 114 Landmesser U, Engberding N, Bahlmann FH, Schaefer A, Wiencke A, Heineke A, Spiekermann S, Hilfiker-Kleiner D, Templin C, Kotlarz D, Mueller M, Fuchs M, Hornig B, Haller H, Drexler H. Statin-induced improvement of endothelial progenitor cell mobilization, myocardial neovascularization, left ventricular function, and survival after experimental myocardial infarction requires endothelial nitric oxide synthase. *Circulation* 2004; 110: 1933-1939 [PMID: 15466656 DOI: 10.1161/01.CIR.0000143232.67642.7A]
- 115 Matsuno H, Takei M, Hayashi H, Nakajima K, Ishisaki A, Kozawa O. Simvastatin enhances the regeneration of endothelial cells via VEGF secretion in injured arteries. *J Cardiovasc Pharmacol* 2004; 43: 333-340 [PMID: 15076215 DOI: 10.1097/00005344-20040300 0-000021
- 116 Liao JK, Shin WS, Lee WY, Clark SL. Oxidized low-density lipoprotein decreases the expression of endothelial nitric oxide synthase. *J Biol Chem* 1995; 270: 319-324 [PMID: 7529227 DOI: 10.1074/jbc.270.1.319]
- 117 Liao JK, Clark SL. Regulation of G-protein alpha i2 subunit expression by oxidized low-density lipoprotein. *J Clin Invest* 1995; 95: 1457-1463 [PMID: 7706449 DOI: 10.1172/JCI117816]
- 118 Tamai O, Matsuoka H, Itabe H, Wada Y, Kohno K, Imaizumi T. Single LDL apheresis improves endothelium-dependent vasodilatation in hypercholesterolemic humans. *Circulation* 1997; 95: 76-82 [PMID: 8994420 DOI: 10.1161/01.CIR.95.1.76]
- 119 López-Farré A, Caramelo C, Esteban A, Alberola ML, Millás I, Montón M, Casado S. Effects of aspirin on platelet-neutrophil interactions. Role of nitric oxide and endothelin-1. *Circulation* 1995; 91: 2080-2088 [PMID: 7895367 DOI: 10.1161/01.CIR.91.7.2080]
- 120 Latini R, Bianchi M, Correale E, Dinarello CA, Fantuzzi G, Fresco C, Maggioni AP, Mengozzi M, Romano S, Shapiro L. Cytokines in

- acute myocardial infarction: selective increase in circulating tumor necrosis factor, its soluble receptor, and interleukin-1 receptor antagonist. *J Cardiovasc Pharmacol* 1994; **23**: 1-6 [PMID: 7511719 DOI: 10.1097/00005344-199401000-00001]
- 121 Ma XL, Weyrich AS, Lefer DJ, Lefer AM. Diminished basal nitric oxide release after myocardial ischemia and reperfusion promotes neutrophil adherence to coronary endothelium. *Circ Res* 1993; 72: 403-412 [PMID: 8418991 DOI: 10.1161/01.RES.72.2.403]
- 122 Assmus B, Urbich C, Aicher A, Hofmann WK, Haendeler J, Rössig L, Spyridopoulos I, Zeiher AM, Dimmeler S. HMG-CoA reductase inhibitors reduce senescence and increase proliferation of endothelial progenitor cells via regulation of cell cycle regulatory genes. *Circ Res* 2003; 92: 1049-1055 [PMID: 12676819 DOI: 10.1161/01.RES.0000070067.64040.7C]
- 123 Kusuyama T, Omura T, Nishiya D, Enomoto S, Matsumoto R, Murata T, Takeuchi K, Yoshikawa J, Yoshiyama M. The effects of HMG-CoA reductase inhibitor on vascular progenitor cells. *J Pharmacol Sci* 2006; 101: 344-349 [PMID: 16891763 DOI: 10.1254/jphs.FP0060102]
- 124 Laufs U, Fata VL, Liao JK. Inhibition of 3-hydroxy-3-methylglutaryl (HMG)-CoA reductase blocks hypoxia-mediated down-regulation of endothelial nitric oxide synthase. *J Biol Chem* 1997; 272: 31725-31729 [PMID: 9395516 DOI: 10.1074/jbc.272.50.31725]
- 125 Davis ME, Cai H, McCann L, Fukai T, Harrison DG. Role of c-Src in regulation of endothelial nitric oxide synthase expression during exercise training. Am J Physiol Heart Circ Physiol 2003; 284: H1449-H1453 [PMID: 12595302 DOI: 10.1152/ ajpheart.00918.2002]
- 126 Kosmidou I, Moore JP, Weber M, Searles CD. Statin treatment and 3' polyadenylation of eNOS mRNA. Arterioscler Thromb Vasc Biol 2007; 27: 2642-2649 [PMID: 17916773 DOI: 10.1161/ ATVBAHA.107.154492]
- 127 Minami Y, Satoh M, Maesawa C, Takahashi Y, Tabuchi T, Itoh T, Nakamura M. Effect of atorvastatin on microRNA 221 / 222 expression in endothelial progenitor cells obtained from patients with coronary artery disease. *Eur J Clin Invest* 2009; 39: 359-367 [PMID: 19371267 DOI: 10.1111/j.1365-2362.2009.02110.x]
- 128 Petersen CP, Bordeleau ME, Pelletier J, Sharp PA. Short RNAs repress translation after initiation in mammalian cells. *Mol Cell* 2006; 21: 533-542 [PMID: 16483934 DOI: 10.1016/j.molcel.2006.01.031]
- 129 Felli N, Fontana L, Pelosi E, Botta R, Bonci D, Facchiano F, Liuzzi F, Lulli V, Morsilli O, Santoro S, Valtieri M, Calin GA, Liu CG, Sorrentino A, Croce CM, Peschle C. MicroRNAs 221 and 222 inhibit normal erythropoiesis and erythroleukemic cell growth via kit receptor down-modulation. *Proc Natl Acad Sci U S A* 2005; 102: 18081-18086 [PMID: 16330772 DOI: 10.1073/pnas.0506216102]
- 130 Cerda A, Fajardo CM, Basso RG, Hirata MH, Hirata RD. Role of microRNAs 221/222 on statin induced nitric oxide release in human endothelial cells. *Arq Bras Cardiol* 2015; 104: 195-201 [PMID: 25517390 DOI: 10.5935/abc.20140192]
- 131 Kuehbacher A, Urbich C, Dimmeler S. Targeting microRNA expression to regulate angiogenesis. *Trends Pharmacol Sci* 2008; 29: 12-15 [PMID: 18068232 DOI: 10.1016/j.tips.2007.10.014]
- 132 Bader AG, Kang S, Zhao L, Vogt PK. Oncogenic PI3K deregulates transcription and translation. *Nat Rev Cancer* 2005; 5: 921-929 [PMID: 16341083 DOI: 10.1038/nrc1753]
- 133 Dimmeler S, Zeiher AM. Akt takes center stage in angiogenesis signaling. Circ Res 2000; 86: 4-5 [PMID: 10625297 DOI: 10.1161/01.RES.86.1.4]
- 134 Whitman M, Downes CP, Keeler M, Keller T, Cantley L. Type I phosphatidylinositol kinase makes a novel inositol phospholipid, phosphatidylinositol-3-phosphate. *Nature* 1988; 332: 644-646 [PMID: 2833705 DOI: 10.1038/332644a0]
- 135 Toker A, Cantley LC. Signalling through the lipid products of phosphoinositide-3-OH kinase. *Nature* 1997; 387: 673-676 [PMID: 9192891 DOI: 10.1038/42648]
- 136 Karar J, Maity A. PI3K/AKT/mTOR Pathway in Angiogenesis. Front Mol Neurosci 2011; 4: 51 [PMID: 22144946 DOI: 10.3389/fnmol.2011.00051]



- 137 Kawasaki K, Smith RS, Hsieh CM, Sun J, Chao J, Liao JK. Activation of the phosphatidylinositol 3-kinase/protein kinase Akt pathway mediates nitric oxide-induced endothelial cell migration and angiogenesis. *Mol Cell Biol* 2003; 23: 5726-5737 [PMID: 12897144 DOI: 10.1128/MCB.23.16.5726-5737.2003]
- 138 Ellis LM, Hicklin DJ. VEGF-targeted therapy: mechanisms of anti-tumour activity. *Nat Rev Cancer* 2008; 8: 579-591 [PMID: 18596824 DOI: 10.1038/nrc2403]
- Morales-Ruiz M, Fulton D, Sowa G, Languino LR, Fujio Y, Walsh K, Sessa WC. Vascular endothelial growth factor-stimulated actin reorganization and migration of endothelial cells is regulated via the serine/threonine kinase Akt. *Circ Res* 2000; 86: 892-896 [PMID: 10785512 DOI: 10.1161/01.RES.86.8.892]
- 140 Papapetropoulos A, García-Cardeña G, Madri JA, Sessa WC. Nitric oxide production contributes to the angiogenic properties of vascular endothelial growth factor in human endothelial cells. *J Clin Invest* 1997; 100: 3131-3139 [PMID: 9399960 DOI: 10.1172/ JCI119868]
- 141 Fulton D, Gratton JP, McCabe TJ, Fontana J, Fujio Y, Walsh K, Franke TF, Papapetropoulos A, Sessa WC. Regulation of endothelium-derived nitric oxide production by the protein kinase Akt. *Nature* 1999; 399: 597-601 [PMID: 10376602 DOI: 10.1038/21218]
- 142 Dimmeler S, Dernbach E, Zeiher AM. Phosphorylation of the endothelial nitric oxide synthase at ser-1177 is required for VEGFinduced endothelial cell migration. FEBS Lett 2000; 477: 258-262 [PMID: 10908731 DOI: 10.1016/S0014-5793(00)01657-4]
- 143 Bahlmann FH, De Groot K, Spandau JM, Landry AL, Hertel B, Duckert T, Boehm SM, Menne J, Haller H, Fliser D. Erythropoietin regulates endothelial progenitor cells. *Blood* 2004; 103: 921-926 [PMID: 14525788 DOI: 10.1182/blood-2003-04-1284]
- 144 Rikitake Y, Liao JK. Rho GTPases, statins, and nitric oxide. Circ Res 2005; 97: 1232-1235 [PMID: 16339495 DOI: 10.1161/01. RES.0000196564.18314.23]
- 145 Lemoine S, Zhu L, Legallois D, Massetti M, Manrique A, Hanouz JL. Atorvastatin-induced cardioprotection of human myocardium is mediated by the inhibition of mitochondrial permeability transition pore opening via tumor necrosis factor-α and Janus kinase/signal transducers and activators of transcription pathway. *Anesthesiology* 2013; 118: 1373-1384 [PMID: 23416384 DOI: 10.1097/ALN.0b013-e31828a7039]
- 146 Sanada S, Asanuma H, Minamino T, Node K, Takashima S, Okuda H, Shinozaki Y, Ogai A, Fujita M, Hirata A, Kim J, Asano Y, Mori H, Tomoike H, Kitamura S, Hori M, Kitakaze M. Optimal windows of statin use for immediate infarct limitation: 5'-nucleotidase as another downstream molecule of phosphatidylinositol 3-kinase. *Circulation* 2004; 110: 2143-2149 [PMID: 15451788 DOI: 10.1161/01.CIR.0000143830.59419.73]
- 147 Solenkova NV, Solodushko V, Cohen MV, Downey JM. Endogenous adenosine protects preconditioned heart during early minutes of reperfusion by activating Akt. *Am J Physiol Heart Circ Physiol* 2006; 290: H441-H449 [PMID: 16155103 DOI: 10.1152/ajpheart.00589.2005]
- 148 Hausenloy DJ, Yellon DM. New directions for protecting the heart against ischaemia-reperfusion injury: targeting the Reperfusion Injury Salvage Kinase (RISK)-pathway. *Cardiovasc Res* 2004; 61: 448-460 [PMID: 14962476 DOI: 10.1016/j.cardiores.2003.09.024]
- 149 Efthymiou CA, Mocanu MM, Yellon DM. Atorvastatin and myocardial reperfusion injury: new pleiotropic effect implicating multiple prosurvival signaling. *J Cardiovasc Pharmacol* 2005; 45: 247-252 [PMID: 15725950 DOI: 10.1097/01.fjc.000-0154376.82445.06]
- 150 Hibbert B, Ma X, Pourdjabbar A, Simard T, Rayner K, Sun J, Chen YX, Filion L, O'Brien ER. Pre-procedural atorvastatin mobilizes endothelial progenitor cells: clues to the salutary effects of statins on healing of stented human arteries. *PLoS One* 2011; 6: e16413 [PMID: 21283543 DOI: 10.1371/journal.pone.0016413]
- 151 Baran Ç, Durdu S, Dalva K, Zaim Ç, Dogan A, Ocakoglu G, Gürman G, Arslan Ö, Akar AR. Effects of preoperative short term use of atorvastatin on endothelial progenitor cells after coronary

- surgery: a randomized, controlled trial. *Stem Cell Rev* 2012; **8**: 963-971 [PMID: 22076751 DOI: 10.1007/s12015-011-9321-z]
- 152 Schmidt-Lucke C, Fichtlscherer S, Rössig L, Kämper U, Dimmeler S. Improvement of endothelial damage and regeneration indexes in patients with coronary artery disease after 4 weeks of statin therapy. Atherosclerosis 2010; 211: 249-254 [PMID: 20211468 DOI: 10.1016/j.atherosclerosis.2010.02.007]
- 153 Tousoulis D, Andreou I, Tsiatas M, Miliou A, Tentolouris C, Siasos G, Papageorgiou N, Papadimitriou CA, Dimopoulos MA, Stefanadis C. Effects of rosuvastatin and allopurinol on circulating endothelial progenitor cells in patients with congestive heart failure: the impact of inflammatory process and oxidative stress. *Atherosclerosis* 2011; 214: 151-157 [PMID: 21122851 DOI: 10.1016/j.atherosclerosis.201 0.11.002]
- 154 Jialal I, Devaraj S, Singh U, Huet BA. Decreased number and impaired functionality of endothelial progenitor cells in subjects with metabolic syndrome: implications for increased cardiovascular risk. Atherosclerosis 2010; 211: 297-302 [PMID: 20171637 DOI: 10.1016/j.atherosclerosis.2010.01.036]
- 155 Cohen MV, Yang XM, Downey JM. Nitric oxide is a preconditioning mimetic and cardioprotectant and is the basis of many available infarct-sparing strategies. *Cardiovasc Res* 2006; 70: 231-239 [PMID: 16376869 DOI: 10.1016/j.cardiores.2005.10.021]
- 156 Scalia R, Gooszen ME, Jones SP, Hoffmeyer M, Rimmer DM, Trocha SD, Huang PL, Smith MB, Lefer AM, Lefer DJ. Simvastatin exerts both anti-inflammatory and cardioprotective effects in apolipoprotein E-deficient mice. *Circulation* 2001; 103: 2598-2603 [PMID: 11382730 DOI: 10.1161/01.CIR.103.21.2598]
- 157 Wassmann S, Laufs U, Müller K, Konkol C, Ahlbory K, Bäumer AT, Linz W, Böhm M, Nickenig G. Cellular antioxidant effects of atorvastatin in vitro and in vivo. *Arterioscler Thromb* Vasc Biol 2002; 22: 300-305 [PMID: 11834532 DOI: 10.1161/ hq0202.104081]
- 158 Mensah K, Mocanu MM, Yellon DM. Failure to protect the myocardium against ischemia/reperfusion injury after chronic atorvastatin treatment is recaptured by acute atorvastatin treatment: a potential role for phosphatase and tensin homolog deleted on chromosome ten? J Am Coll Cardiol 2005; 45: 1287-1291 [PMID: 15837263 DOI: 10.1016/j.jacc.2005.01.021]
- 159 Mihos CG, Pineda AM, Santana O. Cardiovascular effects of statins, beyond lipid-lowering properties. *Pharmacol Res* 2014; 88: 12-19 [PMID: 24631782 DOI: 10.1016/j.phrs.2014.02.009]
- Tousoulis D, Andreou I, Antoniades C, Tentolouris C, Stefanadis C. Role of inflammation and oxidative stress in endothelial progenitor cell function and mobilization: therapeutic implications for cardiovascular diseases. *Atherosclerosis* 2008; 201: 236-247 [PMID: 18599065 DOI: 10.1016/j.atherosclerosis.2008.05.034]
- 161 Yao EH, Yu Y, Fukuda N. Oxidative stress on progenitor and stem cells in cardiovascular diseases. *Curr Pharm Biotechnol* 2006; 7: 101-108 [PMID: 16724944 DOI: 10.2174/138920106776597685]
- Morel DW, DiCorleto PE, Chisolm GM. Endothelial and smooth muscle cells alter low density lipoprotein in vitro by free radical oxidation. *Arteriosclerosis* 1984; 4: 357-364 [PMID: 6466193 DOI: 10.1161/01.ATV.4.4.357]
- 163 Lamb DJ, Wilkins GM, Leake DS. The oxidative modification of low density lipoprotein by human lymphocytes. *Atherosclerosis* 1992; 92: 187-192 [PMID: 1632847 DOI: 10.1016/0021-9150(92)90277-N]
- 164 Lim S, Barter P. Antioxidant effects of statins in the management of cardiometabolic disorders. *J Atheroscler Thromb* 2014; 21: 997-1010 [PMID: 25132378 DOI: 10.5551/jat.24398]
- 165 Carracedo J, Merino A, Briceño C, Soriano S, Buendía P, Calleros L, Rodriguez M, Martín-Malo A, Aljama P, Ramírez R. Carbamylated low-density lipoprotein induces oxidative stress and accelerated senescence in human endothelial progenitor cells. FASEB J 2011; 25: 1314-1322 [PMID: 21228221 DOI: 10.1096/fi.10-173377]
- 166 Tso C, Martinic G, Fan WH, Rogers C, Rye KA, Barter PJ. High-density lipoproteins enhance progenitor-mediated endothelium repair in mice. *Arterioscler Thromb Vasc Biol* 2006; 26: 1144-1149 [PMID: 16528007 DOI: 10.1161/01.ATV.0000216600.37436.cf]



- 167 Griendling KK, FitzGerald GA. Oxidative stress and cardiovascular injury: Part II: animal and human studies. *Circulation* 2003; 108: 2034-2040 [PMID: 14581381 DOI: 10.1161/01. CIR.0000093661.90582.c4]
- 168 Thum T, Fraccarollo D, Schultheiss M, Froese S, Galuppo P, Widder JD, Tsikas D, Ertl G, Bauersachs J. Endothelial nitric oxide synthase uncoupling impairs endothelial progenitor cell mobilization and function in diabetes. *Diabetes* 2007; 56: 666-674 [PMID: 17327434 DOI: 10.2337/db06-0699]
- 169 Urbich C, Knau A, Fichtlscherer S, Walter DH, Brühl T, Potente M, Hofmann WK, de Vos S, Zeiher AM, Dimmeler S. FOXO-dependent expression of the proapoptotic protein Bim: pivotal role for apoptosis signaling in endothelial progenitor cells. FASEB J 2005; 19: 974-976 [PMID: 15824087 DOI: 10.1096/fj.04-2727fje]
- 170 Fleissner F, Thum T. Critical role of the nitric oxide/reactive oxygen species balance in endothelial progenitor dysfunction. Antioxid Redox Signal 2011; 15: 933-948 [PMID: 20712407 DOI: 10.1089/ars.2010.3502]
- 171 Tepper OM, Galiano RD, Capla JM, Kalka C, Gagne PJ, Jacobowitz GR, Levine JP, Gurtner GC. Human endothelial progenitor cells from type II diabetics exhibit impaired proliferation, adhesion, and incorporation into vascular structures. *Circulation* 2002; 106: 2781-2786 [PMID: 12451003 DOI: 10.1161/01. CIR.0000039526.42991.93]
- 172 Tao J, Yang Z, Wang JM, Wang LC, Luo CF, Tang AL, Dong YG, Ma H. Shear stress increases Cu/Zn SOD activity and mRNA expression in human endothelial progenitor cells. *J Hum Hypertens* 2007; 21: 353-358 [PMID: 17287843 DOI: 10.1038/sj.jhh.1002147]
- 173 **He T**, Peterson TE, Holmuhamedov EL, Terzic A, Caplice NM, Oberley LW, Katusic ZS. Human endothelial progenitor cells tolerate oxidative stress due to intrinsically high expression of manganese superoxide dismutase. *Arterioscler Thromb Vasc Biol* 2004; **24**: 2021-2027 [PMID: 15319267 DOI: 10.1161/01. ATV.0000142810.27849.8f]
- 174 Dernbach E, Urbich C, Brandes RP, Hofmann WK, Zeiher AM, Dimmeler S. Antioxidative stress-associated genes in circulating progenitor cells: evidence for enhanced resistance against oxidative stress. *Blood* 2004; 104: 3591-3597 [PMID: 15161665 DOI: 10.1182/blood-2003-12-4103]
- 175 Endres M. Statins: potential new indications in inflammatory conditions. *Atheroscler Suppl* 2006; 7: 31-35 [PMID: 16503422 DOI: 10.1016/j.atherosclerosissup.2006.01.005]
- 176 Bauer SM, Goldstein LJ, Bauer RJ, Chen H, Putt M, Velazquez OC. The bone marrow-derived endothelial progenitor cell response is impaired in delayed wound healing from ischemia. *J Vasc Surg* 2006; 43: 134-141 [PMID: 16414400 DOI: 10.1016/j.jvs.2005.08.038]
- 177 Masuda H, Kalka C, Takahashi T, Yoshida M, Wada M, Kobori M, Itoh R, Iwaguro H, Eguchi M, Iwami Y, Tanaka R, Nakagawa Y, Sugimoto A, Ninomiya S, Hayashi S, Kato S, Asahara T. Estrogenmediated endothelial progenitor cell biology and kinetics for physiological postnatal vasculogenesis. *Circ Res* 2007; 101: 598-606 [PMID: 17656679 DOI: 10.1161/CIRCRESAHA.106.144006]
- 178 Ii M, Nishimura H, Iwakura A, Wecker A, Eaton E, Asahara T, Losordo DW. Endothelial progenitor cells are rapidly recruited to myocardium and mediate protective effect of ischemic preconditioning via "imported" nitric oxide synthase activity. *Circulation* 2005; 111: 1114-1120 [PMID: 15723985 DOI: 10.1161/01.CIR.0000157144. 24888.7E]
- 179 Zhang Y, Devries ME, Skolnick J. Structure modeling of all identified G protein-coupled receptors in the human genome. PLoS Comput Biol 2006; 2: e13 [PMID: 16485037 DOI: 10.1371/journal.

- pcbi.0020013]
- 180 Bjarnadóttir TK, Gloriam DE, Hellstrand SH, Kristiansson H, Fredriksson R, Schiöth HB. Comprehensive repertoire and phylogenetic analysis of the G protein-coupled receptors in human and mouse. *Genomics* 2006; 88: 263-273 [PMID: 16753280 DOI: 10.1016/j.ygeno.2006.04.001]
- 181 Schöneberg T, Hofreiter M, Schulz A, Römpler H. Learning from the past: evolution of GPCR functions. *Trends Pharmacol Sci* 2007; 28: 117-121 [PMID: 17280721 DOI: 10.1016/j.tips.2007.01.001]
- 182 Pierce KL, Premont RT, Lefkowitz RJ. Seven-transmembrane receptors. *Nat Rev Mol Cell Biol* 2002; 3: 639-650 [PMID: 12209124 DOI: 10.1038/nrm908]
- 183 Perez DM. The evolutionarily triumphant G-protein-coupled receptor. *Mol Pharmacol* 2003; 63: 1202-1205 [PMID: 12761327 DOI: 10.1124/mol.63.6.1202]
- 184 Rosenbaum DM, Rasmussen SG, Kobilka BK. The structure and function of G-protein-coupled receptors. *Nature* 2009; 459: 356-363 [PMID: 19458711 DOI: 10.1038/nature08144]
- 185 Venkatakrishnan AJ, Deupi X, Lebon G, Tate CG, Schertler GF, Babu MM. Molecular signatures of G-protein-coupled receptors. *Nature* 2013; 494: 185-194 [PMID: 23407534 DOI: 10.1038/nature11896]
- 186 Callihan P, Mumaw J, Machacek DW, Stice SL, Hooks SB. Regulation of stem cell pluripotency and differentiation by G protein coupled receptors. *Pharmacol Ther* 2011; 129: 290-306 [PMID: 21073897 DOI: 10.1016/j.pharmthera.2010.10.007]
- 187 Kobayashi NR, Hawes SM, Crook JM, Pébay A. G-protein coupled receptors in stem cell self-renewal and differentiation. Stem Cell Rev 2010; 6: 351-366 [PMID: 20625855 DOI: 10.1007/ s12015-010-9167-9]
- 188 Nature Reviews Drug Discovery GPCR Questionnaire Participants. The state of GPCR research in 2004. Nat Rev Drug Discov 2004; 3: 575, 577-626 [PMID: 15272499 DOI: 10.1038/nrd1458]
- 189 Jacoby E, Bouhelal R, Gerspacher M, Seuwen K. The 7 TM G-protein-coupled receptor target family. *ChemMedChem* 2006; 1: 761-782 [PMID: 16902930 DOI: 10.1002/cmdc.200600134]
- 190 Kozasa T, Jiang X, Hart MJ, Sternweis PM, Singer WD, Gilman AG, Bollag G, Sternweis PC. p115 RhoGEF, a GTPase activating protein for Galpha12 and Galpha13. *Science* 1998; 280: 2109-2111 [PMID: 9641915 DOI: 10.1126/science.280.5372.2109]
- 191 Noma K, Oyama N, Liao JK. Physiological role of ROCKs in the cardiovascular system. Am J Physiol Cell Physiol 2006; 290: C661-C668 [PMID: 16469861 DOI: 10.1152/ajpcell.00459.2005]
- 192 Taussig R, Iñiguez-Lluhi JA, Gilman AG. Inhibition of adenylyl cyclase by Gi alpha. *Science* 1993; 261: 218-221 [PMID: 8327893 DOI: 10.1126/science.8327893]
- 193 Morello F, Perino A, Hirsch E. Phosphoinositide 3-kinase signalling in the vascular system. *Cardiovasc Res* 2009; 82: 261-271 [PMID: 19038971 DOI: 10.1093/cvr/cvn325]
- 194 Gilman AG. G proteins: transducers of receptor-generated signals. Annu Rev Biochem 1987; 56: 615-649 [PMID: 3113327 DOI: 10.1146/annurev.bi.56.070187.003151]
- 195 Ghanemi A. Targeting G protein coupled receptor-related pathways as emerging molecular therapies. *Saudi Pharm J* 2015; 23: 115-129 [PMID: 25972730 DOI: 10.1016/j.jsps.2013.07.007]
- 196 Lin SH, Civelli O. Orphan G protein-coupled receptors: targets for new therapeutic interventions. *Ann Med* 2004; 36: 204-214 [PMID: 15181976 DOI: 10.1080/07853890310024668]
- 197 Lynch JR, Wang JY. G Protein-Coupled Receptor Signaling in Stem Cells and Cancer. Int J Mol Sci 2016; 17: [PMID: 27187360 DOI: 10.3390/ijms17050707]
- P- Reviewer: Das UN, Jiang B, Lymperopoulos A S- Editor: Qiu S L- Editor: A E- Editor: Wu HL







Published by Baishideng Publishing Group Inc

8226 Regency Drive, Pleasanton, CA 94588, USA

Telephone: +1-925-223-8242

Fax: +1-925-223-8243

E-mail: bpgoffice@wjgnet.com
Help Desk: http://www.wjgnet.com/esps/helpdesk.aspx
http://www.wjgnet.com

