

The endothelial saga: the past, the present, the future

Dragomir N. Serban · Bernd Nilius · Paul M. Vanhoutte

Received: 11 February 2010 / Accepted: 13 February 2010

© Springer-Verlag 2010

Abstract Endothelium-dependent changes in vasomotor tone, whether evoked by vasoactive agents or physical forces, are recognized as essential for the local hemodynamic control in various normal and pathological circumstances. They are based on a complex signaling network within the vascular wall. In recent years, substantial efforts have been made to analyze how such signals are generated and used in the endothelium-dependent control of vascular smooth muscle. The underlying mechanisms vary with species, age, sex, hormonal status, vascular bed studied, caliber of the blood vessels, triggering stimuli, pre-existing vascular tone, oxidative stress, and pathology. Such aspects and many others will be addressed specifically by the authors contributing to this volume.

Keywords Endothelium · Nitric oxide · EDHF · K_{Ca} · TRP · Oxidative stress

D. N. Serban
Laboratory of Cell Physiology and Pharmacology,
Center for Study and Therapy of Pain,
“Gr. T. Popa” University of Medicine and Pharmacy,
16 Universitatii Str.,
700115 Iasi, Romania

B. Nilius
Laboratory of Ion Channel Research,
Department of Molecular Cell Biology, KU Leuven,
Campus Gasthuisberg, O&N 1, Herestraat 49 bus 802,
3000 Leuven, Belgium

P. M. Vanhoutte (✉)
Department of Pharmacology and Pharmacy,
Li Ka Shing Faculty of Medicine, University of Hong Kong,
Laboratory Block 2/F, 21 Sassoon Road, Pokfulam,
Hong Kong, China
e-mail: vanhoutt@hku.hk

The endothelial saga: the past

The endothelial saga started with Robert Furchgott [18, 61], who demonstrated that endothelial cells play an essential role in the relaxation evoked by acetylcholine in isolated arteries, which is mediated by activation of endothelial muscarinic receptors. His simple pharmacological experiments have revolutionized not only vascular pharmacology and physiology but science in general, as they lead to the discovery of the role of nitric oxide (NO) in biology [61]. Using “sandwich” bioassay preparations (a layering of arterial strips with and without endothelium whereby the contractile responses are measured only in the strip without endothelium), he demonstrated that the endothelium-dependence of the response to acetylcholine is due to the diffusion of a vasodilator substance from the endothelial cells to the vascular smooth muscle cells [18]. Having ruled out prostacyclin, which is produced by endothelial cells [41], he called the unknown mediator “endothelium-derived relaxing factor” (EDRF). The existence of endothelium-dependent responses was rapidly confirmed in different laboratories around the world [see 37]. More sophisticated superfusion-bioassay systems permitted to apply pharmacological inhibitors to either the endothelial cells or the effector vascular smooth muscle cells [e.g., 50]. The biological half-life of EDRF was found to be disappointingly brief (in the order of seconds), making identification by conventional chemical techniques impossible. Early pharmacological studies indicated that endothelial cells can generate several other signals leading to endothelium-dependent relaxations [8]. The latter multiple signals (Fig. 1) eventually became known as “endothelium-derived hyperpolarizing factor(s)” (EDHF), which play a prominent role in smaller arteries and resistance vessels [7, 15]. In addition, it soon became obvious that, in veins [9], and in

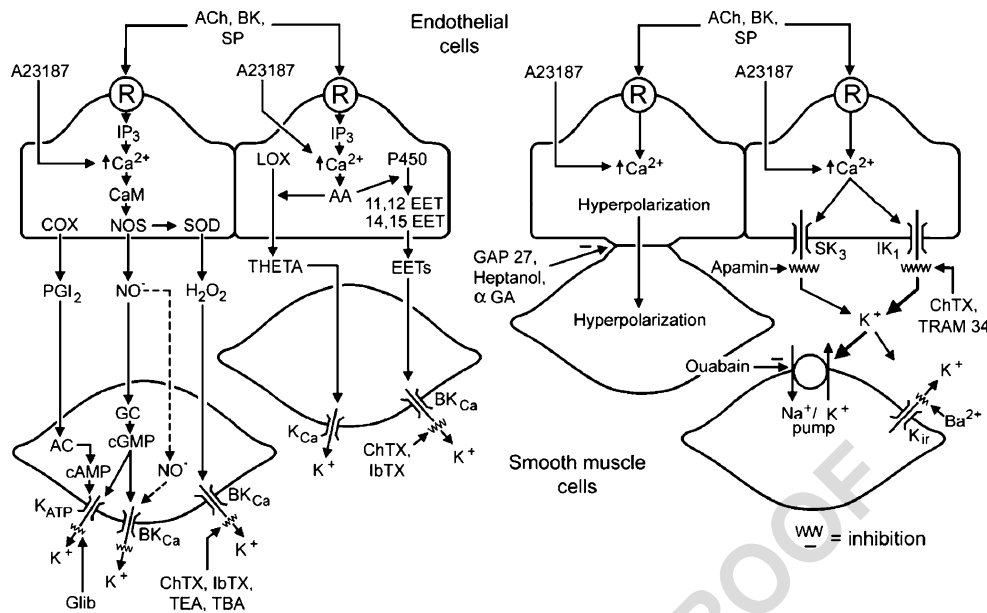


Fig. 1 EDRFs in 2009. Substances such as acetylcholine (*ACh*), bradykinin (*BK*), and substance P (*SP*), through the activation of M_3 -muscarinic, B_2 -bradykinin, and NK_1 -neurokinin receptor subtypes, respectively, and agents that increase intracellular calcium, such as the calcium ionophore A23187, release EDHFs. *CaM* calmodulin, *COX* cyclooxygenase, *EET* epoxyeicosatrienoic acid, *IP₃* inositol triphosphate, *GC* guanylate cyclase, *NAPE* *N*-acylphosphatylethanolamine, *Hyperpol* hyperpolarization, *NOS* NO synthase, O_2^- superoxide anions, *PGI₂* prostacyclin, *P450* cytochrome P450 monooxygenase, *R* receptor, *X* putative EDHF synthase. SR141716 is an antagonist of the cannabinoid receptor subtype CB₁. Glibenclamide (*Glib*) is a selective inhibitor of ATP-sensitive potassium channels (K_{ATP}). Tetraethylammonium (*TEA*) and tetrabutylammonium (*TBA*) are

nonspecific inhibitors of potassium channels when used at high concentrations (>5 mM), while at lower concentrations (1–3 mM), these drugs are selective for calcium-activated potassium channels (K_{Ca}). Iberitoxin (*IBX*) is a specific inhibitor of large conductance K_{Ca} (BK_{Ca}). Charybdotoxin (*CTX*) is an inhibitor of BK_{Ca} , intermediate conductance K_{Ca} (IK_{Ca}), and voltage-dependent potassium channels. Apamin is a specific inhibitor of small conductance K_{Ca} (SK_{Ca}). Barium (Ba^{2+}), in the micromolar range, is a specific inhibitor of the inward rectifier potassium channel (K_{ir}). GAP 27 is an 11-amino acid peptide possessing conserved sequence homology to a portion of the second extracellular loop of connexin. 18 α -glycyrrhetic acid (αGA), and heptanol are gap junction uncouplers (from Vanhoutte et al., 2009. By permission)

59 arteries as well [36], the endothelium produces “endothelium-derived contracting factors” (EDCF), which add to the
 60 difficulty of analyzing endothelium-dependent responses
 61 [17, 37, 64]. More physiological stimuli than acetylcholine
 62 [including physical forces (increases in shear stress), circulating
 63 hormones (catecholamines, vasopressin), platelet products (serotonin, adenosine diphosphate), autacoids (histamine,
 64 bradykinin), prostaglandin E₄, and thrombin] were shown to
 65 cause endothelium-dependent relaxations [37, 63]. Of those
 66 more physiological stimuli, increases in shear stress [51]
 67 explain the endothelium-dependency of flow-mediated
 68 vasodilatation, a response that allows the most accurate
 69 assessment of endothelial function in humans. Research in
 70 the field was fostered by the fact that endothelium-
 71 dependent relaxations are reduced under a number of
 72 pathological conditions, including myocardial infarction
 73 [32] and hypertension [31], which lead to the current
 74 conviction that endothelial dysfunction precedes, or at
 75 least accompanies, vascular disease and predicts the
 76 occurrence of cardiovascular events [39, 63].

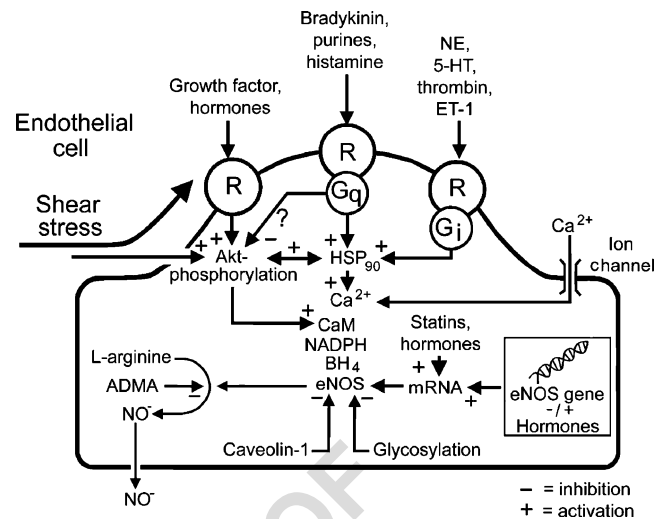
77 It soon appeared that EDRF, whatever its nature,
 78 stimulated soluble guanylyl cyclase in vascular smooth
 79

80 muscle [28]. Soluble guanylyl cyclase catalyzes the
 81 formation of cyclic guanosine monophosphate (cyclic
 82 GMP), which in turn initiates relaxation. The demonstration
 83 followed that under superfusion-bioassay conditions super-
 84 oxide anions scavenge EDRF [21, 52]. Based on the earlier
 85 observation that nitric oxide (NO) activates soluble gua-
 86 nylyl cyclase and is scavenged by superoxide anions [42]
 87 and on his own work with acidified nitrite, Robert
 88 Furchgott proposed in 1986 that his EDRF is NO [19].
 89 Louis Ignarro had reached the same conclusion [27]. One
 90 year later, Salvador Moncada and his colleagues demon-
 91 strated, using a chemiluminescence technique, that, when
 92 cultured endothelial cells are stimulated with bradykinin,
 93 they indeed release NO [45]. The biology of NO was born.
 94 One crucial finding was that macrophages and endothelial
 95 cells transform the semi-essential amino acid L-arginine
 96 into NO and citrulline, and then inhibitors of the respon-
 97 sible enzymatic activity were discovered [24, 46, 48]. The
 98 access to inhibitors of nitric oxide synthase (NOS)
 99 permitted the exploration of the physiological role of NO
 100 in isolated tissues and organs, and in the intact organism.
 101 The fact that they augment arterial blood pressure in vivo
 102

103 [49] implied a role for NO in cardiovascular homeostasis.
 104 The use of NOS inhibitors in vivo rapidly lead to the
 105 conclusion that NO not only is a key player in vasomotor
 106 control but it affects almost every bodily function. The next
 107 breakthrough came when Salomon Snyder and his group
 108 isolated NOS from the brain [e.g., 6]. We now know that
 109 there are three isoforms of the enzyme: neuronal NOS
 110 (nNOS, NOS 1), inducible NOS (iNOS, NOS 2), and
 111 endothelial NOS (eNOS, NOS 3). Paul Huang and
 112 colleagues genetically engineered mice with deletion of
 113 the eNOS gene [25]. These animals have an increased
 114 arterial blood pressure, illustrating the role of NO in the
 115 control of cardiovascular homeostasis.

116 **The endothelial saga: the present**

117 The advent of genetically modified animals and of
 118 inhibitors of NOS permits the systematic exploration of
 119 the role of NO in vascular health and disease, considerably
 120 increasing our knowledge (Fig. 2). In a given blood vessel,
 121 the level of activity of eNOS and the amounts of
 122 endothelium-derived NO released are not constant. They
 123 can be upregulated by chronic increases in shear stress
 124 (exercise), hormones (estrogens), and diet (ω_3 -unsaturated
 125 fatty acids or polyphenols of red wine, green tea, and dark
 126 chocolate). The endothelial production of NO is reduced by
 127 high glucose (diabetes) and increased oxidative stress
 128 (hypertension) [see 3, 63]. NO not only affects the tone of
 129 vascular smooth muscle, but also inhibits platelet aggregation,
 130 in synergy with endothelium-derived prostacyclin [47], and
 131 the growth of the media [55]. It reduces the endothelial
 132 production of endothelin-1 [62] and of cyclooxygenase-derived
 133 EDCF [14]. NO inhibits the expression of endothelial adhesion
 134 molecules and, thus, the adhesion of platelets and white blood
 135 cells [47, 63]. It modulates angiogenesis [3, 68]. The
 136 signaling cascade, in particular, the role of Akt, in the
 137 phosphorylation that leads to activation of eNOS is unraveled
 138 [3, 16, 29, 33]. The original concept that the eNOS is a
 139 strictly Ca^{2+} -dependent enzyme, and, thus, that endothelium-
 140 dependent relaxations rely entirely on an increase in intracellular
 141 Ca^{2+} -concentration, has been challenged [3, 16]. Moreover,
 142 in vivo responses to acetylcholine in arterioles consist of two
 143 phases: (a) a rapidly conducted vasodilatation initiated by
 144 a local rise in endothelial Ca^{2+} but independent of
 145 endothelial Ca^{2+} -signaling at remote sites and (b) a slower
 146 complementary dilatation associated with a Ca^{2+} -wave that
 147 propagates along the endothelium [57]. In the mouse aorta,
 148 calcium-imaging shows that only some clusters of endothelial
 149 cells respond to acetylcholine, which represent only one
 150 third of the total number of cells, but this is enough for
 151 endothelium-dependent relaxation [4]. The importance of
 152



153 **Fig. 2** Possible mechanisms by which production of nitric oxide is
 154 regulated in endothelial cells. Nitric oxide is produced through
 155 enzymatic conversion of L-arginine by NOS (endothelial or type III,
 156 eNOS). The transcription of this enzyme is regulated genomically
 157 by hormones and growth factors. Stability of eNOS mRNA is
 158 modulated by statins and hormones. eNOS enzyme activity
 159 requires calcium, calmodulin, nicotinamide adenine dinucleotide
 160 phosphate (NADPH), and 5, 6, 7, 8-tetra-hydrobiopterine
 161 (BH₄). Enzyme activity is regulated by complexing to these
 162 proteins in microdomains of the endothelial cell. Association
 163 with this complex of heat shock protein 90 (HSP₉₀) increases
 164 enzyme activity. Stimulation of specific receptors on the
 165 endothelial surface (R) complexed with guanine nucleotide
 166 regulatory proteins, which are sensitive to pertussis toxin
 167 (G_i) or insensitive to pertussis toxin (G_q), activate
 168 intracellular pathways that modulate eNOS activity posttrans-
 169 lationally through heat shock protein 90 or Akt-phosphorylation.
 170 Association of eNOS with caveolin-1 or glycosylation of the
 171 enzyme reduces activity. A metabolite of L-arginine, asym-
 172 metric dimethyl arginine (ADMA) decreases production of
 173 the nitric oxide through competitive binding to eNOS. Thus,
 174 this endogenous amine may be a risk factor for the develop-
 175 ment of cardiovascular disease. *Plus signs* indicate stimula-
 176 tion, *minus signs* indicate inhibition, *question marks* indicate
 177 those pathways in which the regulation is unknown (from Van-
 178 houette et al., 2009. By permission)

153 the caveolae for the activity of eNOS is now established
 154 [20, 40]. The formation of NO-metabolites constitutes a
 155 non-enzymatic source of activators of soluble guanylyl
 156 cyclase [38]. Beyond NO itself, derivatives such as nitroxyl
 157 (HNO) and nitrosothiols have also emerged as EDRFs and
 158 HNO may be as important as NO in rodent small arteries
 159 [2]. The binding of NO to superoxide anions, with the
 160 formation of peroxynitrite, is a major player in genesis of
 161 endothelial dysfunction [23, 30, 58, 67]. The progressive
 162 inability of endothelial cells, prematurely aged by the
 163 exposure to risk factors, to generate sufficient NO may
 164 well be the initial step permitting the inflammatory
 165 response that leads to atherosclerosis [see 63]. The most
 166 widely used therapeutic agents for the treatment of
 167 cardiovascular disease enhance the ability of the endothelial
 168 cells to produce NO [26, 59, 64].

169 We now appreciate better the importance and the
 170 complexity of endothelium-dependent hyperpolarization in
 171 the local control of vascular tone [7]. Although EDHF has
 172 been considered to be of particular importance in smaller
 173 arteries, we have to recognize that its contribution to
 174 vasodilatation may be merely transient [22]. Nevertheless,
 175 coordinated increases in small artery diameter occur by
 176 means of flow-mediated vasodilatation (shear-stress-in-
 177 duced and NO-dependent) combined with the conducted
 178 vasodilatation resulting from electrotonic propagation of
 179 hyperpolarization in the endothelium [56]. At the level of
 180 endothelial protrusions, functional cooperation ensures the
 181 EDHF-component of endothelium-dependent vasodilata-
 182 tion, which is mediated by K^+ released from endothelium
 183 and involves endothelial $K_{Ca2.3}$ and $K_{Ca3.1}$, local intersti-
 184 tial Ca^{2+} , Ca^{2+} -sensing receptors co-localized with $K_{Ca3.1}$
 185 in caveolin-poor regions of endothelial cells, myo-
 186 endothelial gap junctions, and the Na/K pump and $K_{ir2.1}$
 187 of the vascular smooth muscle [11]. Experiments in
 188 dysgenic mice suggest that $K_{Ca2.3}$ and $K_{Ca3.1}$ have
 189 important but different contributions to endothelium-
 190 dependent vasodilatation and, thus, represent novel thera-
 191 peutic targets for the treatment of hypertension [5, 66].
 192 Inositol 1,4,5-endothelial trisphosphate receptors in the
 193 endothelial protrusions subserve local Ca^{2+} -release events
 194 (“pulsars”), which activate the functionally co-localized
 195 $K_{Ca3.1}$ [34]. Activators of the small and intermediate
 196 conductance K channels constitute useful pharmacological
 197 tools and potential new drugs for the treatment of
 198 hypertension [54].

199 The Ca^{2+} -dependent component of local vasodilation
 200 obviously depends on Ca^{2+} influx into endothelial cells.
 201 One of the most attractive candidate influx pathways has
 202 been the store-operated Ca^{2+} entry (SOC), which could be
 203 mediated by transient receptor potential (TRP) channels
 204 [see 43 for a critical review]. SOC was indeed identified in
 205 endothelium [1, 13], but a direct relation to NO produc-
 206 tion and release is still under evaluation [4]. Non-store-
 207 operated channels seem to play a more important role
 208 in regulation of NO release [65]. The involvement of
 209 TRPV4-channels in flow-induced endothelium-dependent
 210 vasodilatation is now generally accepted [35, see also 44
 211 for a review]; the mechanism requires an active CYP
 212 epoxygenase and channel translocation to the cell mem-
 213 brane, where it is associated with caveolin-1. Moreover,
 214 the expression of caveolin-1 is required for EDHF-related
 215 relaxation, by modulating the membrane location and
 216 activity of TRPV4 channels and connexins, which are
 217 both implicated at different steps in the EDHF-signaling
 218 pathway [53]. The TRPV4 channels of both endothelial
 219 and vascular smooth muscle cells are critically involved in
 220 endothelium-dependent vasodilatation of mesenteric arter-
 221 ies and in TRPV4-knockout mice the hypertension

induced by NOS inhibition is greater than in wild-type
 animals [12].

The endothelial saga: the future

Much remains to be learned about the precise regulation of
 NO release by endothelial cells and also about the
 consequences of its perturbation within the complex chain
 of events leading to the vascular dysfunction characteristic of
 hypertension, diabetes, and atherosclerosis [64]. We still do
 not completely understand the exact role of EDHF-mediated
 responses in physiology and pathology, as we are still unable
 to selectively interfere with them in vivo [7]. We still do not
 fully comprehend the importance of EDCFs in endothelial
 dysfunction [60]. Finally, we have to unravel the complex
 interactions between the different endothelium-derived sig-
 nals. For example, in diabetic mice, hyperglycemia-induced
 changes in endothelial function are linked to COX2 and
 oxidative stress (enhanced NADPH oxidase and decreased
 SOD expression), uncoupling of eNOS, and changes in its
 expression and regulation, while EDHF-mediated vasodila-
 tion can be maintained, but with a modified profile [10].
 Whatever the future of endothelial research will yield, we
 should not forget that this extraordinary scientific saga
 started with the very simple pharmacological experiments
 of Robert Furchgott [18, 61], whose memory we honor in
 this special issue.

Acknowledgement This work was supported by grants from the
 Romanian National Authority for Scientific Research via UEFISCSU
 and CNCSIS (grant ID_1156/2007-2010 from the program IDEI of
 plan PNCDI-II), the Belgian Ministry for Science Policy (Interuniver-
 sity Attraction Pole IUAP P6/28), the Research Foundation-Flanders
 (G.0172.03 and G.0565.07), the Research Council of the KU Leuven
 (GOA 2004/07 and EF/95/010).

The editors initiated this special issue as a result of discussions
 originating at “Works and Views in Endothelium-Dependent Vasodi-
 lation,” an international symposium organized by D. N. Serban in
 May 2009, in Iasi, Romania, which was supported from the mentioned
 Romanian grant and sponsored by DABMMed.

References

1. Abdullaev IF, Bisaillon JM, Potier M, Gonzalez JC, Motiani RK, Trebak M (2009) Stim1 and Orai1 mediate CRAC currents and store-operated calcium entry important for endothelial cell proliferation. *Circ Res* 103:1289–1299
2. Andrews KL, Irvine JC, Tare M, Apostolopoulos J, Favalaro JL, Triggle CR, Kemp-Harper BK (2009) A role for nitroxyl (HNO) as an endothelium-derived relaxing and hyperpolarizing factor in resistance arteries. *Br J Pharmacol* 157:540–550
3. Balligand J-L, Feron O, Dessy C (2009) eNOS activation by physical forces: from short-term regulation of contraction to chronic remodeling of cardiovascular tissues. *Physiol Rev* 80:481–534

273	4. Boittin FX, Gribi F, Serir K, Bény JL (2008) Ca ²⁺ -independent PLA2 controls endothelial store-operated Ca ²⁺ entry and vascular tone in intact aorta. <i>Am J Physiol Heart Circ Physiol</i> 295:H2466–H2474	339
274		340
275		341
276		342
277	5. Brähler S, Kaistha A, Schmidt VJ, Wöfle SE, Busch C, Kaistha BP, Kacic M, Hasenau AL, Grgic I, Si H, Bond CT, Adelman JP, Wulff H, de Wit C, Hoyer J, Köhler R (2009) Genetic deficit of SK3 and IK1 channels disrupts the endothelium-derived hyperpolarizing factor vasodilator pathway and causes hypertension. <i>Circulation</i> 119:2323–2332	343
278		344
279		345
280		346
281		347
282		348
283	6. Bredt DS, Snyder SH (1990) Isolation of nitric oxide synthetase, a calmodulin-requiring enzyme. <i>Proc Nat Acad Sci U S A</i> 87:682–685	349
284		350
285		351
286	7. Busse R, Edwards G, Félétou M, Fleming I, Vanhoutte PM (2002) EDHF: Bringing the concepts together. <i>Trends Pharmacol Sci</i> 23:374–380	352
287		353
288		354
289	8. De Mey JG, Claeys M, Vanhoutte PM (1982) Endothelium-dependent inhibitory effects of acetylcholine, adenosine triphosphate, thrombin and arachidonic acid in the canine femoral artery. <i>J Pharmacol Exp Ther</i> 222:166–173	355
290		356
291		357
292		358
293	9. De Mey JG, Vanhoutte PM (1982) Heterogeneous behavior of the canine arterial and venous wall: importance of the endothelium. <i>Circ Res</i> 51:439–447	359
294		360
295		361
296		362
297	10. Ding H, Aljofan M, Triggle CR (2007) Oxidative stress and increased eNOS and NADPH oxidase expression in mouse microvessel endothelial cells. <i>J Cell Physiol</i> 212:682–689	363
298		364
299	11. Dora KA, Gallagher NT, McNeish A, Garland CJ (2008) Modulation of endothelial cell KCa _{3.1} channels during endothelium-derived hyperpolarizing factor signaling in mesenteric resistance arteries. <i>Circ Res</i> 102:1247–1255	365
300		366
301		367
302		368
303	12. Earley S, Pauyo T, Drapp R, Tavares MJ, Liedtke W, Brayden JE (2009) TRPV4-dependent dilation of peripheral resistance arteries influences arterial pressure. <i>Am J Physiol Heart Circ Physiol</i> . doi:10.1152/ajpheart.00241.2009	369
304		370
305		371
306		372
307	13. Fasolato C, Nilius B (1998) Store depletion triggers the calcium release-activated calcium current (ICRAC) in macrovascular endothelial cells: a comparison with Jurkat and embryonic kidney cell lines. <i>Pflügers Arch</i> 436:69–74	373
308		374
309		375
310		376
311	14. Feletou M, Tang EH, Vanhoutte PM (2008) Nitric oxide the gatekeeper of endothelial vasomotor control. <i>Front Biosci</i> 13:4198–4217	377
312		378
313		379
314	15. Félétou M, Vanhoutte PM (2006) EDHF: where are we now? <i>Arterioscler Thromb Vasc Biol</i> 26:1215–1225	380
315		381
316	16. Fleming I, Busse R (2003) Molecular mechanisms involved in the regulation of the endothelial nitric oxide synthase. <i>Am J Physiol Regul Integr Comp Physiol</i> 284:R1–R12	382
317		383
318		384
319	17. Furchgott RF, Vanhoutte PM (1989) Endothelium-derived relaxing and contracting factors. <i>FASEB J</i> 3:2007–2017	385
320		386
321	18. Furchgott RF, Zawadzki JV (1980) The obligatory role of endothelial cells in the relaxation of arterial smooth muscle by acetylcholine. <i>Nature</i> 299:373–376	387
322		388
323		389
324	19. Furchgott RF (1988) Studies on relaxation of rabbit aorta by sodium nitrite: the basis for the proposal that acid-activable inhibitory factor from bovine retractor penis is inorganic nitrite and the endothelium-derived relaxing factor is nitric oxide. In: Vanhoutte PM (ed) <i>Vasodilatation: vascular smooth muscle peptides, autonomic nerves and endothelium</i> . Raven, New York, pp 401–414	390
325		391
326		392
327		393
328		394
329		395
330		396
331	20. Gratton JP, Bernatchez P, Sessa WC (2004) Caveolae and caveolins in the cardiovascular system. <i>Circ Res</i> 94:1408–1417	397
332		398
333		399
334	21. Gryglewski RJ, Palmer RMJ, Moncada S (1986) Superoxide anion is involved in the breakdown of endothelium-derived vascular relaxing factor. <i>Nature</i> 320:454–456	400
335		401
336		402
337	22. Harrington LS, Carrier MJ, Gallagher N, Gilroy D, Garland CJ, Mitchell JA (2007) Elucidation of the temporal relationship between endothelial-derived NO and EDHF in mesenteric vessels. <i>Am J Physiol Heart Circ Physiol</i> 293:H1682–H1688	403
338		404
	23. Heistad DD, Wakisaka Y, Miller J, Chu Y, Pena-Silva R (2009) Novel aspects of oxidative stress in cardiovascular diseases. <i>Circ J</i> 73:201–207	
	24. Hibbs JB Jr, Taintor RR, Vavrin Z (1987) Macrophage cytotoxicity: role for L-arginine deiminase and imino nitrogen oxidation to nitrite. <i>Science</i> 235:473–476	
	25. Huang PL, Huang Z, Mashimo H, Bloch KD, Moskowitz MA, Bevan JA, Fishman MC (1995) Hypertension in mice lacking the gene for endothelial nitric oxide synthase. <i>Nature</i> 377:239–242	
	26. Ignarro LJ (2008) Different pharmacological properties of two enantiomers in a unique β -blocker, nebivolol. <i>Cardiovasc Ther</i> 26:115–134	
	27. Ignarro LJ, Byrns RE, Wood KS (1988) Biochemical and pharmacological properties of endothelium-derived relaxing factor and its similarity to nitric oxide radical. In: Vanhoutte PM (ed) <i>Vasodilatation: vascular smooth muscle peptides, autonomic nerves and endothelium</i> . Raven, New York, pp 427–436	
	28. Ignarro LJ, Harbison RG, Wood KS, Kadowitz PJ (1986) Activation of purified soluble guanylate cyclase by endothelium-derived relaxing factor from intrapulmonary artery and vein: stimulation by acetylcholine, bradykinin and arachidonic acid. <i>J Pharmacol Exp Ther</i> 237:893–900	
	29. Jagannadan D, Sessa WC, Fulton D (2005) Intracellular location regulates calcium-calmodulin-dependent activation of organelle-restricted eNOS. <i>Am J Physiol Cell Physiol</i> 289:C1024–C1033	
	30. Kojda G, Harrison D (1999) Interactions between NO and reactive oxygen species: pathophysiological importance in atherosclerosis, hypertension, diabetes and heart failure. <i>Cardiovasc Res</i> 43:562–571	
	31. Konishi M, Su C (1983) Role of endothelium in dilator responses of spontaneously hypertensive rat arteries. <i>Hypertension</i> 5:881–886	
	32. Ku DD (1982) Coronary vascular reactivity after acute myocardial infarction. <i>Science</i> 218:576–578	
	33. Lamas S, Lowenstein CJ, Michel T (2007) Nitric oxide signaling comes of age: 20 years and thriving. <i>Cardiovasc Res</i> 75:207–209	
	34. Ledoux J, Taylor MS, Bonev AD, Hannah RM, Solodushko V, Shui B, Tallini Y, Kotlikoff MI, Nelson MT (2008) Functional architecture of inositol 1, 4, 5-trisphosphate signaling in restricted spaces of myoendothelial projections. <i>Proc Natl Acad Sci U S A</i> 105:9627–9632	
	35. Loot AE, Popp R, Fisslthaler B, Vriens J, Nilius B, Fleming I (2008) Role of cytochrome P450-dependent transient receptor potential V4 activation in flow-induced vasodilatation. <i>Cardiovasc Res</i> 80:445–452	
	36. Lüscher TF, Vanhoutte PM (1986) Endothelium-dependent contractions to acetylcholine in the aorta of the spontaneously hypertensive rat. <i>Hypertension</i> 8:344–348	
	37. Lüscher TF, Vanhoutte PM (1990) The endothelium: modulator of cardiovascular function. CRC, Boca Raton	
	38. Lundberg JO (2006) Nitric oxide metabolites and cardiovascular disease. Markers, mediators, or both? <i>J Am Col Cardiol</i> 47:580–581	
	39. Marin E, Sessa WC (2007) Role of endothelial-derived nitric oxide in hypertension and renal diseases. <i>Curr Opin Nephrol Hypertens</i> 16:105–110	
	40. Minshall RD, Sessa WC, Stan RV, Anderson RG, Malik AB (2003) Caveolin regulation of endothelial function. <i>Am J Physiol Lung Cell Mol Physiol</i> 285:L1179–L1183	
	41. Moncada S, Vane JR (1979) Pharmacology and endogenous roles of prostaglandin endoperoxides, thromboxane A ₂ and prostacyclin. <i>Pharmacol Rev</i> 30:293–331	
	42. Murad F, Mitta CK, Arnold WP, Katsuki S, Kimura H (1978) Guanylate cyclase: activation by azide, nitro-compounds, nitric	

405 oxide and hydroxyl radical an inhibition by haemoglobin and 447
 406 myoglobin. *Adv Cycl Nucleotide Res* 9:145–158 448
 407 43. Nilius B, Droogmans G, Wondergem R (2003) Transient receptor 449
 408 potential channels in endothelium: solving the calcium entry 450
 409 puzzle? *Endothelium* 10:5–15 451
 410 44. Nilius B, Owsianik G, Voets T, Peters JA (2007) Transient receptor 452
 411 potential cation channels in disease. *Physiol Rev* 87:165–217 453
 412 45. Palmer RMJ, Ashton DS, Moncada S (1988) Vascular endothelial 454
 413 cells synthesize nitric oxide from L-arginine. *Nature* 333:664–666 455
 414 46. Palmer RMJ, Moncada S (1989) A novel citrulline-forming 456
 415 enzyme implicated in the formation of nitric oxide by vascular 457
 416 endothelial cells. *Biochem Biophys Res Commun* 158:348–352 458
 417 47. Radomski MW, Palmer RMJ, Moncada S (1987) The role of nitric 459
 418 oxide and cGMP in platelet adhesion to vascular endothelium. 460
 419 *Biochem Biophys Res Commun* 148:1482–1489 461
 420 48. Rees DD, Palmer RMJ, Hodson HF, Moncada S (1989) A specific 462
 421 inhibitor of nitric oxide formation from L-arginine attenuates 463
 422 endothelium-dependent relaxation. *Br J Pharmacol* 96:418–424 464
 423 49. Rees DD, Palmer RMJ, Moncada S (1989) The role of 465
 424 endothelium-derived nitric oxide in the regulation of blood 466
 425 pressure. *Proc Natl Acad Sci U S A* 86:3375–3378 467
 426 50. Rubanyi GM, Lorenz RR, Vanhoutte PM (1985) Bioassay of 468
 427 endothelium-derived relaxing factor(s). Inactivation by catechol- 469
 428 amines. *Am J Physiol* 249:H95–H101 470
 429 51. Rubanyi GM, Romero JC, Vanhoutte PM (1986) Flow-induced 471
 430 release of endothelium-derived relaxing factor. *Am J Physiol* 250: 472
 431 H1145–H1149 473
 432 52. Rubanyi GM, Vanhoutte PM (1986) Superoxide anions and 474
 433 hyperoxia inactivate endothelium-derived relaxing factor(s). *Am* 475
 434 *J Physiol* 250:H822–H827 476
 435 53. Saliez J, Bouzin C, Rath G, Ghisdal P, Desjardins F, Rezzani R, 477
 436 Rodella LF, Vriens J, Nilius B, Feron O, Balligand JL, Dessy C 478
 437 (2008) Role of caveolar compartmentation in endothelium-derived 479
 438 hyperpolarizing factor-mediated relaxation: Ca²⁺ signals and gap 480
 439 junction function are regulated by caveolin in endothelial cells. 481
 440 *Circulation* 117:1065–1074 482
 441 54. Sankaranarayanan A, Raman G, Busch C, Schultz T, Zimin PI, 483
 442 Hoyer J, Köhler R, Wulff H (2009) Naphtho[1, 2-d]thiazol-2- 484
 443 ylamine (SKA-31), a new activator of KCa₂ and KCa_{3.1} 485
 444 potassium channels, potentiates the endothelium-derived hyper- 486
 445 polarizing factor response and lowers blood pressure. *Mol* 487
 446 *Pharmacol* 75:281–295 488
 447 55. Scott-Burden T, Vanhoutte PM (1993) The endothelium as a 447
 448 regulator of vascular smooth muscle proliferation. *Circulation* 87: 448
 449 V51–V55 449
 450 56. Segal SS (2005) Regulation of blood flow in the microcirculation. 450
 451 *Microcirculation* 12:33–45 451
 452 57. Tallini YN, Brekke JF, Shui B, Doran R, Hwang SM, Nakai J, 452
 453 Salama G, Segal SS, Kotlikoff MI (2007) Propagated endothelial 453
 454 Ca²⁺ waves and arteriolar dilation in vivo: measurements in 454
 455 Cx40BAC GCaMP2 transgenic mice. *Circ Res* 101:1300–1309 455
 456 58. Touyz RM (2004) Reactive oxygen species and angiotensin II 456
 457 signaling in vascular cells— implications in cardiovascular 457
 458 disease. *Braz J Med Biol Res* 37:1263–1273 458
 459 59. Vanhoutte PM (1998) Endothelial dysfunction and inhibition of 459
 460 converting enzyme. *Eur Heart J* 19:J7–J15 460
 461 60. Vanhoutte PM (2010) COX-1 and vascular disease. *Clin Pharma-* 461
 462 *col Ther* (in press) 462
 463 61. Vanhoutte PM (2009) How we learned to say NO. *Arterioscler* 463
 464 *Thromb Vasc Biol* 29:1156–1160 464
 465 62. Vanhoutte PM (2009) Say NO to ET. *J Auton Nerv Syst* 81:271– 465
 466 277 466
 467 63. Vanhoutte PM, Shimokawa H, Tang EH, Feletou M (2009) 467
 468 Endothelial dysfunction and vascular disease. *Acta Physiol* 468
 469 196:193–222 469
 470 64. Vanhoutte PM, Tang EH (2008) Endothelium-dependent contrac- 470
 471 tions: when a good guy turns bad! *J Physiol* 586:5295–5304 471
 472 65. Vriens J, Owsianik G, Fisslthaler B, Suzuki M, Janssens A, Voets 472
 473 T, Morisseau C, Hammock BD, Fleming I, Busse R, Nilius B 473
 474 (2005) Modulation of the Ca²⁺ permeable cation channel TRPV4 474
 475 by cytochrome P450 epoxygenases in vascular endothelium. *Circ* 475
 476 *Res* 97:908–915 476
 477 66. Wölflle SE, Schmidt VJ, Hoyer J, Köhler R, de Wit C (2009) 477
 478 Prominent role of KCa_{3.1} in endothelium-derived hyperpolarizing 478
 479 factor-type dilations and conducted responses in the microcircu- 479
 480 lation in vivo. *Cardiovasc Res* 82:476–483 480
 481 67. Wolin MS (2009) Reactive oxygen species and the control of 481
 482 vascular function. *Am J Physiol Heart Circ Physiol* 296:H539– 482
 483 H549 483
 484 68. Yu J, deMuinck ED, Zhuang Z, Drinane M, Kauser K, Rubanyi 484
 485 GM, Qian HS, Murata T, Escalante B, Sessa WC (2005) 485
 486 Endothelial nitric oxide synthase is critical for ischemic remodel- 486
 487 ling, mural cell recruitment, and blood flow reserve. *Proc Natl* 487
 488 *Acad Sci U S A* 102:10999–11004 488

Q2

AUTHOR QUERIES

AUTHOR PLEASE ANSWER ALL QUERIES.

- Q1. Please check if the authors' affiliations were presented correctly.
- Q2. Please check if the publication data of reference item number 60 need to be updated.

UNCORRECTED PROOF