Heterogeneity and emergent behaviour in the vascular endothelium
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The endothelium is the single layer of cells lining all blood vessels, and it is a remarkable cardiovascular control centre. Each endothelial cell has only a small number (on average six) of interconnected neighbours. Yet this arrangement produces a large repertoire of behaviours, capable of controlling numerous cardiovascular functions in a flexible and dynamic way. The endothelium regulates the delivery of nutrients and removal of waste by regulating blood flow and vascular permeability. The endothelium regulates blood clotting, responses to infection and inflammation, the formation of new blood vessels, and remodelling of the blood vessel wall. To carry out these roles, the endothelium autonomously interprets a complex environment crammed with signals from hormones, neurotransmitters, pericytes, smooth muscle cells, various blood cells, viral or bacterial infection and proinflammatory cytokines. It is generally assumed that the endothelium responds to these instructions with coordinated responses in a homogeneous population of endothelial cells. Here, we highlight evidence that shows that neighbouring endothelial cells are highly heterogeneous and display different sensitivities to various activators. Cells with various sensitivities process different extracellular signals into distinct streams of information in parallel, like a vast switchboard. Communication occurs among cells and new ‘emergent’ signals are generated that are non-linear composites of the inputs. Emergent signals cannot be predicted or deduced from the properties of individual cells. Heterogeneity and emergent behaviour bestow capabilities on the endothelial collective that far exceed those of individual cells. The implications of heterogeneity and emergent behaviour for understanding vascular disease and drug discovery are discussed.

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Introduction
The endothelium regulates virtually every cardiovascular activity by releasing numerous vasoactive agents. To regulate blood flow and blood pressure, endothelium-derived nitric oxide and prostacyclins promote vasodilation, whereas production of endothelin, superoxide and thromboxane stimulate vasoconstriction [1]. Blood fluidity is regulated by endothelial factors that interrupt the blood clotting cascade (antithrombin III, the protein C receptor thrombomodulin, and tissue factor pathway inhibitor [2]), platelet activation (nitric oxide and prostacyclin, exonucleotidases and surface heparan sulphates [2]) and fibrinolysis (tissue-type plasminogen activator and its inhibitor plasminogen activator inhibitor-1 [3]). Angiogenesis is regulated by the endothelium receiving signals from molecules such as vascular endothelial growth factor (VEGF) and von Willebrand factor [4-8]. Blood vessel permeability plays a key role in tissue perfusion and host defence and is regulated by the expression of endothelial cell adhesion molecules. Endothelial cells also play a key role in immune and inflammatory reactions by regulating leukocyte movement into tissues via production of chemokines, colony stimulating factors and expression of specific proteins and cell adhesion molecules at sites requiring defence or repair [9-11].

In making ‘decisions’ on vasoactive outputs, the endothelium must coordinate responses across many cells to numerous local and circulating cues. The cues are generated by changes in physiological status and contained in signals within the chemical environment to which the endothelium is exposed. The environment is complex. Signals are received in the form of the blood composition (e.g. pH, O_2), hormones, neurotransmitters, and from pericytes, smooth muscle cells, various blood cells, viral or bacterial infection, proinflammatory cytokines and even endothelial cells themselves. Multiple signals may arrive simultaneously, each providing separate instructions to the vascular system [12-16]. To manage and accurately process the information, the endothelium must selectively and sensitively resolve each message held within the chemical environment while remaining resistant to the random spontaneous activity in cells (noise).

Even a single message may itself vary widely in intensity and carry different instructions at different concentrations. Major physiological crisis events, such as bleeding, may generate large local concentrations of activators [17]
to evoke substantial immediate cardiovascular responses, for example, blood clotting. However, a great deal of cardiovascular activity is routine, small background adjustments driven by modest changes in circulating activators. Fluctuations in the concentration of agents that activate the endothelium to produce background physiological adjustments are vanishingly small and require the endothelium to be exceptionally sensitive. The concentration changes of circulating hormones, such as leptin, estradiol, parathyroid hormone, epinephrine, angiotensin II are small and vary periodically from a basal value of a few tens of picomolar to a peak rise in the low hundreds of picomolar [12-15,18]. Leptin, for example, varies periodically over a 24-hour period from a basal concentration of ~49 nM to a peak of 99 nM (8-16 ng/ml) [12]. Estradiol oscillates between 74 nM (20 pg/ml) and 220 nM (60 pg/ml) though occasionally reached an unusually high concentration of 1.5 nM (400 pg/ml) [12]. Parathyroid hormone from 5 pM (50 ng/l) to 80 pM (75 ng/l) [13] and epinephrine from 275 pM (50 pg/ml) to 2.2 nM (400 pg/ml) [19]. Even during a fight-or-flight crisis epinephrine concentration may reach only ~55 nM (10 ng/ml) [20]. Each of these concentrations would be considered very low by conventional pharmacological standards. Yet the high-sensitivity enables the endothelium to effectively respond and detect signals from circulating activators to maintain ‘background’ cardiovascular activity.

However, a problem coupled to the high-sensitivity of the endothelium is how random ‘noise’ occurring in the detection system can be rejected. Random noise is unavoidable and arises from stochastic processes that occur in the cells themselves. Random noise can evoke signals of a substantial magnitude. The endothelium must recognise these events as noise and prevent them from evoking widespread activity while retaining high-sensitivity. These features of endothelial sensitivity raise a number of questions. How does the endothelium efficiently detect exceptionally low concentrations of activators associated with background physiological events, while also being able to respond to very high concentrations during crisis (e.g. bleeding) events? How is noise distinguished from ‘real’ signal? By what mechanism does the endothelium process multiple, separate, and even contradictory, messages? What underlies the change in endothelial sensitivity that occurs in cardiovascular disease? The answers to these questions are largely unknown but will provide fundamental insight into endothelial function and the dysfunction occurring in cardiovascular disease.

In this review, we outline key properties of the endothelium that are involved in the detection of the numerous signals required for cardiovascular activity. We highlight how the behaviour of individual endothelial cells differs from expectations derived from the most common experimental approaches used to study the endothelium and the impact these differences may have on drug discovery.

Uniformly heterogenous
Given the differences in haemodynamics, physiology and structure of blood vessels, it is not surprising that endothelial cell specialisation occurs in different parts of the vascular system [21-33,34]. However, within a region of a blood vessel, most studies examining sensing and activation treat the endothelium as a homogenous population of cells that respond uniformly. Each cell, it is believed, detects each signal equally, and each cell’s response is considered to be a miniature version of the entire response. An attractive feature of the proposal is that no additional consideration is required to explain coordinated function of the endothelium; coordination is achieved by uniform activation of cells.

The hypothesis that endothelial cells behave uniformly is implicit in most experimental approaches used to study the endothelium. In organ bath experiments, the mechanical response of the artery or vein is used as an indirect measure of endothelial activity. This type of experiment has yielded many important insights into endothelial function (e.g. discovery of nitric oxide) but averages the behaviour of thousands of endothelial cells and assumes each cell behaves in a similar way. The organ bath approach is not unique in assuming uniformity. Many current interpretations of gene expression, protein levels or metabolic signalling that are derived from immuno-blotts, PCR or microarrays also assume that all cells of a population are comparable in receptor complement and signalling processes [35,36]. When changes are reported in cardiovascular disease, it is assumed all cells are altered equally.

While the hypothesis of homogenous cell responses is attractive and widely accepted, the hypothesis has not been confirmed at the level of individual cells. When individual cells in a population have been examined, differences in the time course of responses or expression of proteins have been almost universally reported [37-40]. For example, signalling in cells occurs out of phase with neighbouring regions and receptors are heterogeneously distributed [21,41-47]. The distribution of angiotensin II immunostaining is irregular in neighbouring endothelial cells of femoral mesenteric artery [30]. An uneven mosaic pattern of von Willebrand factor (VWF)-positive and von Willebrand factor (VWF)-negative endothelial cells occurs in many vascular beds and even in capillaries [30,48,49]. Adrenergic α-adrenoceptor clusters and cannabinoid receptor distribution are heterogeneous among cells in the endothelium [42]. There is also heterogeneity in the distribution of muscarinic (M3) receptors and purinergic (P2Y2) receptors in neighbouring cells (Figure 1; [50]).
Functional data also show heterogeneity in endothelial responses. Acetylcholine-evoked Ca^{2+} responses are larger at branches in rat thoracic aorta than that of neighbouring non-branch regions [51]. The reverse is true of histamine [51]. Studies of murine thoracic aorta endothelial cells that used only single concentrations of agonists found that while most cells (82%) responded to ATP, large fractions of cells did not respond to acetylcholine, bradykinin or substance P [52]. Responses to mechanical activation also are not uniform and certain populations of cells are more likely to respond to shear stress [53].

Our results also demonstrate heterogeneity in the endothelium’s response. Different cells respond to different concentrations of the activator acetylcholine [54]. As the concentration increased, an increasing number of cells were recruited. The increasing recruitment of cells was part of the concentration-dependence of the response. While the sensitivity range of the endothelium as a whole to acetylcholine extended over three orders of concentration magnitude, each cell was sensitive over only one order of magnitude [54], that is, the system has properties that differ and exceed the properties of individual parts (cells). This arrangement (having different cells sensitive to various concentrations) solves a problem common to sensory systems, that is, how to create a system that is exceptionally sensitive to a stimulus but does not saturate at low-stimulus intensity. The endothelium’s organisation solves this problem by having individual cells that are highly sensitive detectors of very limited concentration ranges [54].

Interestingly, cells that were comparably sensitive were positioned close together in discrete clusters. As the concentration of each agonist increased, more cells in the cluster activated and more clusters responded [50,54]. These findings revealed that sensing cells are neither randomly nor uniformly distributed, but structured into sensing regions [54].

The clustering of cells creates properties of the system that are absent in single cells and generates a mechanism to help reject noise. Clustering may provide a co-occurrence detection system [50,54]. When single cells in the endothelium were activated (as would occur in stochastic noise events), there is little propagation of signals occurred to neighbouring cells. However, when two or
more neighbouring cells activated together, signal propagation occurred. By rejecting noise, signal detection will improve and randomly occurring events are unlikely to be propagated. These observations again show that the endothelium as a whole has properties that are quite distinct from those of individual cells. Clustering may offer other advantages. Clustering may allow the uptake and breakdown mechanisms for diffusible messengers (e.g. nitric oxide, prostaglandin) to be overwhelmed, providing increased spillover of signals. Clustering may also limit interference from neighbouring cells that are responding to a different stimulus, that is, a single cell responding in isolation may easily be influenced by neighbouring cells and have its signal overridden. A cluster of cells each performing the same task may be much harder to override.

The mechanisms giving rise to the organisation of cells into clusters are not yet clear. Perhaps self-replication occurs during development or the cells may be at different developmental ages because of cellular renewal. Alternatively there may be feedback control of function and receptor expression based on location or a result of a self-organisation process at the cellular level.

Fractured but whole; communication and cooperation

While the system appears to be fractured and operating as a series of distinct clusters of cells, communication ensures the endothelium functions seamlessly as a harmonised whole. Each endothelial cell is connected to approximately six neighbouring cells (Figure 1; [50*]) and interaction occurs among connected cells. Communication via gap junctions [55,56] is an acknowledged feature of the endothelium [55,57–59], and the low electrical resistance (~5–70 MΩ [60–64]) demonstrates the high extent of connectivity among cells.

The communication pathways create a network capable of relaying information [55,57–59,65]. However, the communication pathways in the endothelium are usually treated as being like a wire in a telecommunication system. An input to the system is relayed from cell to cell but decays passively with distance, as described by the systems ‘cable properties’ [66]. The network itself is not thought to interact with the input but relays signals without changing the signals characteristics. In other systems with similar passive network properties, separate input signals may passively summate or cancel in an approximately linear way. In this regard, the system has linear features with ‘resultant’ properties which can be predicted. The resultant may be a sum or a difference of the interacting elements.

However, in contrast to the simple resultant properties of a system, an incredible feature of the endothelium is that the system may interact with the inputs to generate new distinctive signals that differ in a non-linear way from the inputs. The muscarinic agonist carbachol and purinergic agonist ATP, activate spatially separate clusters of cells [50*]. In those separate clusters of cells, Ca²⁺ signals evoked by carbachol and ATP are distinctive [50*]. Carbachol evokes oscillating Ca²⁺ signals, ATP evokes an initial Ca²⁺ spike followed by a decline towards basal values (Figure 2). This observation demonstrates that an input/output relationship exists for each agonist that is temporally as well as spatially distinctive, that is, the endothelium can distinguish inputs and assign outputs (Figure 2).

While most endothelial cells are heterogeneous and particularly sensitive to one activator, a small number of cells are sensitive to more than one activator [50*]. When the agonists are added separately, these cells also generated distinctive signals to each agonist and the signals were similar to those seen in cells sensitive to only one activator (Figure 2; [50*]). This observation suggests that the distinctive signals evoked by each agonist are a feature of the agonist acting on the cells rather than the cell itself.

Emergent signals. Surprisingly, even though separate cells are activated by muscarinic and purinergic agonists, when carbachol and ATP were present together, the resulting Ca²⁺ signal is distinct from those signals generated by either of the agonists in isolation (Figure 2; [50*]).

The change in the response occurring, when both agonists were present, is particularly interesting. The change in the steady-state response is expansive in that it far exceeds either the average or summation of the two inputs (Figure 2; [50*]). The generation of a distinct new signal suggests that cells perform signalling computations and combine information from multiple sources. The change in signal characteristics occur even though distinct, separate cells were activated by each agonist, that is, communication occurs between the separate clusters (Figure 2; [50*]). The computations on the signals show the endothelium is flexible and interacts with the inputs to generate new signals.

The precise nature of the computation carried out on the signals and the underlying mechanisms are not clear. Nonetheless, the computation is important in that it is a feature that emerges from the collective dynamics of the endothelial network and provides a mechanism for the endothelium to interactively monitor external environments via distributed sensing across separate cells. Examining the endothelial system as a whole reveals properties that are greater than the sum of its parts.

Emergent properties: why organise with heterogeneity?

The interactive collective activity of the cells may hold the key to resolving some of the incredible virtues of the
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Figure 2

(a) CCh EC25 First Responding  ATP EC25 First Responding  Combined First Responding

(b) CCh EC25  ATP EC25  Combined

(c) Cell 1  Cell 2  Cell 3

(d) Peak Ca²⁺ Response (Average ΔF/F₀)

(e) Average Ca²⁺ Response 30s after initial Peak (ΔF/F₀)

Unique signals in separate cells. (a) Composite Ca²⁺ images showing cells that respond in the first 4 s of activation by the EC25 concentrations of CCh (left, green) and ATP (middle, red). Images are of the same field of endothelium. The right-panel shows cells activated (cyan) in the same field of endothelium when both drugs were applied together. (b) Ca²⁺ responses from all activate cells in the field of endothelium shown in (a). The Ca²⁺ increase evoked by CCh was (on average) a slow increase that remained elevated on which repeating oscillations occurred (left green). The response to ATP on average was a sharp transient increase that declined towards resting values (red middle). When both agonists were applied (combined; blue right) the Ca²⁺ increase appeared to have features of each agonist, that is, a slow but larger initial increase than with CCh, which remained more elevated than ATP and slowly declined. Agonists were present for the duration indicated by the line above each trace. (c) Examples of responses from three separate cells to CCh, ATP and to the two agonists when applied together. In each panel in (c), traces are from the cells indicated by the white dots in the panels in (a). It is the same three cells in each case. Cell 1 is shown in the left panel in (c), cell 2 in the middle panel and cell 3 in the right panel. Cell 1 (left panel) responds to CCh but not to ATP. The characteristics of the response in Cell 1 is altered when both ATP and CCh are present (combined) with a faster and larger upstroke. Cell 2 responds to ATP but not CCh. Once again, the
endothelium (Figure 3). Endothelial heterogeneity permits different cells to acquire, in an exceptionally sensitive manner, different elements of the overall information available. However, while cells detect a very limited aspect of the total information content, information is shared so that, collectively, the endothelium interprets the entire chemical environment. As a collective, the endothelium has properties that exceed the capabilities of single cells. This feature is not unique to the endothelium. Biological systems are recognised increasingly as having properties that are distinct from the individual components of the system. New distinct features often arise from interactions and give rise to behaviours that are absent when cells are examined in isolation.

New features that appear in a system and which differ from the expectations of the sum of the components are called ‘emergent properties’ [67,68]. Emergent properties differ significantly from those observed in linear (resultant) systems. Unlike linear systems where the whole is equal to the sum of the parts, emergent systems arise from non-linear interactions and create new collective behaviours that make the whole much greater than the sum of the parts. Emergent properties cannot be reduced to properties of the constituent parts of the system and resist attempts at being inferred or predicted by calculation. In the case of the endothelium, distinctive signals are generated to different agonists, and signals appear in the system that are quite different from the sum of the inputs when multiple agonists are present [50*,54*]. The difference between the behaviour of individual cells and the population average result in the endothelium as a whole being capable of processing more information, more precisely, than cells acting alone. These features have important implications for basic investigations on endothelial function and on drug discovery.

Emergent behaviour in cardiovascular disease. The underlying rationale of most basic investigations is the desire to understand the components that give rise to a response. In disease, these components identify targetable processes, for example, protein kinases, receptors or ion channels, for small molecule design. Most drug discovery investigations rely on averages from population (cells or tissue) studies. However, this reliance assumes uniform behaviour in populations, and believes an understanding of how each cell works will result in an understanding of the system, that is, the investigations are driven by reductionism.

In the reductionist approach, a larger system is analysed by breaking it down into pieces and determining the connections between the parts [69]. Isolated cells, proteins, molecules and ions have sufficient explanatory power to provide an understanding of the whole system and the changes that occur in disease (Figure 3). Reductionism has provided a wealth of knowledge on cellular components and their function. However, despite the success of reductionism, it is increasingly clear that biological function can only rarely be attributed to isolated components.

Reductionist approaches have generated insights into how cells work within a system; however, drug discovery requires an understanding of how the overall system works [70]. Yet, drug discovery is largely based on reductionist approaches such as genomics, proteomics, metabolomics, high-throughput screening, combinatorial chemistry and bioinformatics [70]. These approaches have not brought the new products that were anticipated [71,72]. For example, knowledge of the genome sequences of humans and various pathogenic agents, once hailed as the step change opening doors to new drug development, and ‘personalised’ or ‘precision’ medicine, has led to the identification of only a limited number of beneficial drug targets [73]. Gene therapy, stem-cell research, antisense technology and cancer vaccines have not materialised to the degree so feverishly predicted by early supporters [72]. Despite eye-watering financial investment, sixty years of reductionist approaches involving molecular targeting of drugs for specific enzymes and receptors in cancer chemotherapies, have failure rates of 90% in those agents managing to reach end-stage trials [74*].

For cardiovascular drug discovery, an understanding of the behaviour of the endothelium requires an appreciation of the multiple non-linear interactions and feedbacks that occur within and among cells. The interactions and feedbacks are currently poorly understood. Unfortunately, even if a near complete understanding existed, the behaviour of the system would remain difficult to

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(Figure 2 Legend Continued) characteristics of the response in Cell 2 is altered when both ATP and CCh are present (combined) with a more sustained later Ca++ change. Cell 3 responds to each agonist (CCh and to ATP). Once again, the characteristics of the response in Cell 3 is altered when both ATP and CCh are present (combined). (d) Mean peak responses (black circles) to the EC50 of CCh and ATP separately and when both were present together (combined). The red line shows the calculated mean of peak response when both agonists were added separately. The blue shaded region shows the standard error of the mean. The blue line shows the sum of the peak responses when both agonists were added separately. The blue shaded region shows the standard error of the mean. The combined peak response exceeded the mean and was less than the summed response. (e) Mean steady-state responses (black circles) to the EC50 of CCh and ATP separately and when both were present together (combined). The red line shows the calculated mean of the steady-state response when both agonists were added separately. The red shaded region shows the standard error of the mean. The blue line shows the sum of the steady-state responses when both agonists were added separately. The blue shaded region shows the standard error of the mean. The combined steady-state response exceeded both the mean and the summed response. All Scale bars = 50 μm. From Ref. [50*] with permission.
Uniformity, heterogeneity and reductionism. (a) Illustration of a homogenous population of cells responding to single stimuli. Each cell responds uniformly to either activator (green or red) and generates one output to each stimulus. (b) Heterogeneous populations of cells responding to one (bi and i) and two (biii) different stimuli. Separate spatially distinct clusters of cells process and respond to each activator to generate specific outputs. When both activators are present together (biii) the separate regions respond and multiple outputs can be generated. This arrangement permits the endothelium to simultaneously process different stimuli in parallel. (c) Illustration showing a reductionist approach to drug discovery. The normal, steady-state behaviour present in health (ci) may be disrupted in disease (cii). The endothelium may compensate for this alteration by upregulating proteins in other cells to restore a near normal steady-state in disease (ciii). A reductionist approach to drug discovery, that measures the individual components, may attribute this upregulation to the dysfunction in disease rather than the compensatory mechanism employed by the endothelium to overcome the disease. Targeting this upregulated protein may force the endothelium into another new steady-state that is not beneficial (civ) which lacks the compensatory mechanism (cv) present before pharmacological intervention.
predict and reconstruct. Altering the function of a component in the system will have rippling unintentional consequences, because of feedbacks and interactions, that may be difficult to foresee and may rarely be beneficial (Figure 3). Key to understanding the endothelium is one defining principal – the endothelium exists in a complex steady-state. The system resists changes and will continuously work back (asymptotically) towards the steady-state value.

In a cardiovascular disease, alterations of key components (enzymes, ion channels) may trigger a change which forces the entire system into a new steady-state, albeit one that is dysfunctional. However, the new dysfunctional condition will once again be maintained in a steady-state by multiple altered interactions and feedbacks occurring among enzymes, metabolic processes, ion channels and so on. As a result, there are multiple changes in the function of the components of the system (e.g., enzymes, ion channels). Most of the changes will be consequences rather than causes of cardiovascular disease and are, in fact, beneficial acting to stabilise the system and halt or limit the progression of the disease (Figure 3). The triggering, initiating, event may not be easy to identify among the changes. Therefore, targeting pathways blindly, as is done through reductionism (e.g., identified with proteomics), may have adverse effects as stabilising ‘beneficial’ changes will almost certainly be targeted. It may be fruitless to attach any particular significance to changes in specific biomarkers/proteins (e.g., ion channels, enzymes) in cardiovascular disease — they may be consequences rather than causes of the disease. Pharmacologically altering the behaviour of a biomarker/protein may not restore the system. Instead, these pharmacological agents may cause additional changes in overall function of the system and yet another new steady-state could arise. The interactions and feedbacks that occur in complex systems may explain why the development of many, perhaps most, successful clinically used drugs has been through serendipity rather than rational drug design [75]. A successful approach for rational therapeutic development in any system with emergent properties will require an understanding of the multiple interactions among vital components that support the entire network’s structure and function, and how the interactions change in cardiovascular disease.

Conflict of interest statement
Nothing declared.

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References and recommended reading
Papers of particular interest, published within the period of review, have been highlighted as:

● of special interest

1. Sandoo A, van Zanten JJ, Metsios GS, Carroll D, Kitas GD: The endothelium and its role in regulating vascular tone. Open Cardiovasc Med J 2010, 4:302-312.

2. van Hinsbergh VW: Endothelium—role in regulation of coagulation and inflammation. Semin Immunopathol 2012, 34:93-106.

3. Chapin JC, Hajjar KA: Fibrinolysis and the control of blood coagulation. Blood Rev 2015, 29:17-24.

4. Semenza GL: Vascular responses to hypoxia and ischemia. Arterioscler Thromb Vasc Biol 2010, 30:648-652.

5. Frais P, Mazzone M, Schmidt T, Carmeliet P: Regulation of angiogenesis by oxygen and metabolism. Dev Cell 2009, 16:167-179.

6. Maharaj AS, Saint-Geniez M, Maldonado AE, D’Amore PA: Vascular endothelial growth factor localization in the adult. Am J Pathol 2006, 168:639-648.

7. dela Paz NG, Walshe TE, Leach LL, Saint-Geniez M, D’Amore PA: Role of shear-stress-induced VEGF expression in endothelial cell survival. J Cell Sci 2012, 125:831-843.

8. Randi AM, Smith KE, Castaman G: von Willebrand factor regulation of blood vessel formation. Blood 2018, 132:132-140.

9. Poer JS, Sessa WC: Evolving functions of endothelial cells in inflammation. Nat Rev Immunol 2007, 7:803-815.

10. Oynebraten I, Bakke O, Brandtzaeg P, Johansen FE, Haraldsen G: Rapid chemokine secretion from endothelial cells originates from 2 distinct compartments. Blood 2004, 104:314-320.

11. Rajavashisth TB, Andalibi A, Territo MC, Berliner JA, Navab M, Fogelman AM, Luiss AJ: Induction of endothelial cell expression of granulocyte and macrophage colony-stimulating factors by modified low-density lipoproteins. Nature 1990, 344:254-257.

12. Licinio J, Negrao AB, Mantzoros C, Kaklamani V, Wong ML, Bongiorno PB, Mulla A, Cearnal L, Veldhuis JD, Flier JS et al.: Synchronicity of frequently sampled, 24-h concentrations of circulating leptin, luteinising hormone, and estradiol in healthy women. Proc Natl Acad Sci U S A 1998, 95:2541-2546.

13. Kitamura N, Shigeno C, Shiomi K, Lee KC, Ohta S, Sone T, Katsushina S, Tadamura E, Kousaka T, Yamamoto I et al.: Episodic fluctuation in serum intact parathyroid-hormone concentration in men. J Clin Endocrinol Metab 1990, 70:252-263.

14. Schoff C, Becker C, Prank K, Von Zur Muhlen A, Brabant G: Twenty-four-hour rhythms of plasma catecholamines and their relation to cardiovascular parameters in healthy young men. Eur J Endocrinol 1997, 137:675-683.

15. Dott C, Breckling U, Deraid I, Fehm HL, Born J: Plasma epinephrine and norepinephrine concentrations of healthy humans associated with nighttime sleep and morning arousal. Hypertension 1997, 30:71-76.

16. Macmillan D, McCarron JG: Regulation by FK506 and rapamycin of Ca2+ release from the sarcoplasmic reticulum in vascular smooth muscle: the role of FK506 binding proteins and mTOR. Br J Pharmacol 2009, 158:1112-1120.

17. Hubbell JA, McIntire LV: Platelet active concentration profiles near growing thrombi. A mathematical consideration. Biophys J 1988, 55:937-945.

18. Kala R, Fyhruquist F, Eisalo A: Diurnal variation of plasma angiotensin II in man. Scand J Clin Lab Invest 1973, 31:383-385.

19. Genter P, Berman N, Jacob M, Ipp E: Counterregulatory hormones oscillate during steady-state hypoglycemia. Am J Physiol 1998, 275:E821-E829.

20. Wortsman J: Role of epinephrine in acute stress. Endocrinol Metab Clin North Am 2002, 31:79-106.
Interactive signalling in the endothelium McCarron et al. 31

21. Kaur H, Carvalho J, Looso M, Singh P, Chennupati R, Preussner J, Gunther S, Albarara-Juarez J, Tischner D, Classen S et al.: Single-cell profiling reveals heterogeneity and functional patterning of GPCR expression in the vascular system. Nat Commun 2017, 8.

22. Aird WC: Phenotypic heterogeneity of the endothelium: I. Structure, function, and mechanisms. Circ Res 2007, 100:158-173.

23. Stevens T, Phan S, Frid MG, Alvarez D, Herzog E, Stenmark KR: Lung vascular cell heterogeneity: endothelium, smooth muscle, and fibroblasts. Proc Am Thorac Soc 2008, 5:783-791.

24. Reese TS, Kamovsky MJ: Fine structural localization of a blood-brain barrier to exogenous peroxidase. J Cell Biol 1967, 34:207-217.

25. Mehta D, Malik AB: Signaling mechanisms regulating endothelial permeability. Physiol Rev 2006, 86:279-387.

26. Wisse E: An electron microscopic study of the fenestrated endothelial lining of rat liver sinusoids. J Ultrastruct Res 1970, 31:125-150.

27. Vanhoutte PM, Miller VM: Heterogeneity of endothelium-dependent responses in mammalian blood vessels. J Cardiovasc Pharmacol 1985, 7(Suppl. 3):S12-23.

28. Turner RR, Backstead JH, Warrne RA, Wood GS: Endothelial cell phenotypic diversity. In situ demonstration of immunologic and enzymatic heterogeneity that correlates with specific morphologic subtypes. Am J Clin Pathol 1987, 87:569-575.

29. Page C, Rose M, Yacoub M, Pigott R: Antigenic heterogeneity of vascular endothelium. Am J Pathol 1992, 141:673-683.

30. Tomlinson A, Van Vlijmen H, Loesch A, Burnstock G: An immunohistochemical study of cell endothelial heterogeneity in the rat: observations in "en face" Hauthen preparations. Cell Tissue Res 1991, 262:173-181.

31. Lincoln J, Loesch A, Burnstock G: Localization of vasopressin, serotonin and angiotensin II in endothelial cells of the renal and mesenteric arteries of the rat. Cell Tissue Res 1990, 259:341-344.

32. Yamamoto K, de Waard V, Feurs C, Loskutoff DJ: Tissue distribution and regulation of murine von Willebrand factor gene expression in vivo. Blood 1992, 92:2791-2801.

33. Aird WC, Edelberg JM, Weiler-Guettler H, Simmons WW, Smith TW, Rosenberger DG: Vascular bed-specific expression of an endothelial cell gene is programmed by the tissue microenvironment. J Cell Biol 1997, 138:1117-1124.

34. McCarron JG, Lee MD, Wilson C: The endothelium solves problems that endothelial cells do not know exist. Trends Pharmacol Sci 2017, 38:322-339.

This paper described that cooperativity and heterogeneous behaviour along with the network properties of the endothelium bestows a rapid adaptive problem-solving functionality.

35. Regard JB, Sato IT, Coughlin SR: Anatomical profiling of G protein-coupled receptor expression. Cell 2008, 135:561-571.

36. Hakak Y, Shrestha D, Goeggel MC, Behan DP, Chalmers DT: Global analysis of G protein-coupled receptor signaling in human tissues. FEBS Lett 2003, 550:11-17.

37. Tay S, Hughey JJ, Lee TK, Lipniacki T, Quake SR, Covert MW: Single-cell NF-kappaB dynamics reveal digital activation and analogue information processing. Nature 2010, 466:267-271.

38. Birtwistle MR, Rauch J, Kiyatkin A, Aksamiitei E, Dobrzensky M, Hoek JB, Kolch W, Ogunaake BA, Kholodenko BN: Emergence of bimodal cell population responses from the interplay between analog single-cell signaling and protein expression noise. BMC Syst Biol 2012, 6:109.

39. Buettner F, Nataraajan KN, Casale FP, Prospero V, Scialdone A, Theis FJ, Teichmann SA, Marioni JC, Stegle O: Computational analysis of cell-to-cell heterogeneity in single-cell RNA-sequencing data reveals hidden subpopulations of cells. Nat Biotechnol 2015, 33:155-160.

40. Grun D, Lyubimova A, Kester L, Wiebrands K, Basak O, Sasaki N, Clevers H, van Oudenaarden A: Single-cell messenger RNA sequencing reveals rare intestinal cell types. Nature 2015, 525:251-255.

41. Jacobsen JC, Aalkjaer C, Matchkov VV, Nilsson H, Freiberg JJ, Holstein-Rathlou NH: Heterogeneity and weak coupling may explain the synchronisation characteristics of cells in the arterial wall. Philos Trans A Math Phys Eng Sci 2008, 366:3483-3502.

42. Daly CJ, Ross RA, Whyte J, Henstridge CM, Irving AJ, McGrath JC: Fluorescent ligand binding reveals heterogeneous distribution of adrenoceptors and 'cannabinoid-like' receptors in small arteries. Br J Pharmacol 2010, 159:787-796.

43. Cleaver O, Melton DA: Endothelial signaling during development. Nat Med 2003, 9:661-668.

44. Thurliey K, Tovey SC, Moenke G, Prince VL, Meena A, Thomas AP, Skupin A, Taylor CW, Falcke M: Reliable encoding of stimulus intensities within random sequences of intracranial Ca2+ spikes. Sci Signal 2014, 7:ra59.

45. Garland CJ, Bagher P, Powell C, Ye X, Lemmye HAL, Borysova L, Dora KA: Voltage-dependent Ca2+ entry into smooth muscle during contraction promotes endothelium-mediated feedback vasodilation in arterioles. Sci Signal 2017, 10.

46. Francis M, Waldrup JR, Qian X, Solodushko V, Meriwether J, Taylor MS: Functional tuning of intrinsic endothelial Ca2+ dynamics in swine coronary arteries. Circ Res 2016, 118:1078-1090.

47. Taylor MS, Francis M: Decoding dynamic Ca2+ signaling in the vascular endothelium. Front Physiol 2014, 5:447.

48. Yuan L, Chan GC, Beeler D, Janes L, Spokes KC, Dharaneeswaran H, Mojiri A, Adams WJ, Sciuto T, Garcia-Cardena G et al.: A role of stochastic phenotype switching in generating mosaic endothelial cell heterogeneity. Nat Commun 2016, 7:10160.

This paper shows phenotypic heterogeneity exists in the expression of VWF between neighbouring endothelial cells that had been exposed to precisely the same extracellular environment. The difference in expression involved random transitions in the DNA methylation status of the VWF promoter.

49. Senis YA, Richardson M, Tinlin S, Maurice DH, Giles AR: Changes in the pattern of distribution of von Willebrand factor in rat aortic endothelial cells following thrombin generation in vivo. Br J Haematol 1996, 93:193-203.

50. Lee MD, Wilson C, Saunter CD, Kennedy C, Girkin JM, McCarron JG: Spatially structured cell populations process multiple sensory signals in parallel in intact vascular endothelium. Sci Signal 2018, 11.

By simultaneously analysing the responses of hundreds of endothelial cells, this paper was first to show that different activators evoked different signals in the endothelium and that signals combined in a non-linear way when more than one activator was present.

51. Huang TY, Chu TF, Chen HI, Jen CJ: Heterogeneity of [Ca2+]i signaling in intact rat aortic endothelium. FASEB J 2000, 14:797-804.

This paper analysed the individual responses of hundreds of endothelial cells so show pronounced differences in the sensitivity of neighbouring cells to acetylcholine histamine-sensitive. Sensitivity differed by two orders of magnitude in individual cells.

52. Marie I, Beny JL: Calcium imaging of murine thoracic aorta endothelium by confocal microscopy reveals inhomogeneous distribution of endothelial cells responding to vasodilator agents. J Vasc Res 2002, 39:250-257.

53. Wilson C, Lee M, McCarron JG: Acetylcholine released by endothelial cells facilitates flow-mediated dilatation. J Physiol 2016, 594:7267-7307.

54. Wilson C, Saunter CD, Girkin JM, McCarron JG: Clusters of specialized detector cells provide sensitive and high fidelity receptor signaling in intact endothelium. FASEB J 2016, 30:2000-2013.

This paper analysed Ca2+ signals from hundreds of cells while inside pressurised arteries. The paper was showed that cells that were
comparably sensitive to an activator were clustered in small regions of the endothelium to provide a coincidence detection system to reject noise.

55. Socha MJ, Domeier TL, Behringer EJ, Segal SS: Coordination of intercellular Ca\(^{2+}\) signaling in endothelial cell tubes of mouse resistance arteries. Microcirculation 2012, 19:575-770.

56. Emerson GG, Segal SS: Endothelial cell pathway for conduction of hyperpolarization and vasodilation along hamster feed artery. Circ Res 2000, 86:94-100.

57. Socha MJ, Behringer EJ, Segal SS: Calcium and electrical signalling along endothelium of the resistance vasculature. Basic Clin Pharmacol Toxicol 2012, 110:80-86.

58. Tallini YN, Brekke JF, Shui B, Doran R, Hwang SM, Nakai J, Salama G, Segal SS, Kotlikoff MI: Propagated endothelial Ca\(^{2+}\) waves and arteriolar dilation in vivo: measurements inCx40BAC GCaMP2 transgenic mice. Circ Res 2007, 101:1300-1309.

59. Longden TA, Dabertrand F, Koide M, Gonzales AL, Tykocki NR, Brayden JE, Hill-Eubanks D, Nelson MT: Capillary K\(^{+}\)-sensing initiates retrograde hyperpolarization to increase local cerebral blood flow. Nat Neurosci 2017, 20:717-72.

60. Carter TD, Ogden D: Acetylcholine-stimulated changes of membrane potential and intracellular Ca\(^{2+}\) concentration recorded in endothelial cells in situ in the isolated rat aorta. Pflugers Arch 1994, 428:476-484.

61. Marchenko SM, Sage SO: Electrical properties of resting and acetylcholine-stimulated endothelium in intact rat aorta. J Physiol 1993, 462:735-751.

62. Yamamoto Y, Klemm MF, Edwards FR, Suzuki H: Intercellular electrical communication among smooth muscle and endothelial cells in guinea-pig mesenteric arterioles. J Physiol 2001, 535:181-195.

63. Behringer EJ, Socha MJ, Polo-Parada L, Segal SS: Electrical conduction along endothelial cell tubes from mouse feed arteries: confounding actions of glycyrrhetinic acid derivatives. Br J Pharmacol 2012, 166:774-787.

64. Olschewski A, Olschewski H, Brau ME, Hempelmann G, Vogel W, Safronov BV: Basic electrical properties of in situ endothelial cells of small pulmonary arteries during postnatal development. Am J Respir Cell Mol Biol 2001, 25:285-290.

65. Kamishima T, McCarron JG: Ca\(^{2+}\) removal mechanisms in rat cerebral resistance size arteries. Biophys J 1998, 75:1767-1773.

66. Behringer EJ, Segal SS: Tuning electrical conduction along endothelial tubes of resistance arteries through Ca\(^{2+}\)-activated K\(^{+}\) channels. Circ Res 2012, 110:1311-1321.

67. Kacic S: Systems biology, emergence and antireductionism. Saud J Biol Sci 2016, 23:584-591.

68. Van Regenmortel MH: Reductionism and complexity in molecular biology. Scientists now have the tools to unravel biological and overcome the limitations of reductionism. EMBO Rep 2004, 5:1016-1020.

69. Ahn AC, Tewari M, Poon CS, Phillips RS: The limits of reductionism in medicine: could systems biology offer an alternative? PLoS Med 2006, 3:e208.

70. Butcher EC: Can cell systems biology rescue drug discovery? Nat Rev Drug Discov 2005, 4:461-467.

71. Kubinyi H: Drug research: myths, hype and reality. Nat Rev Drug Discov 2003, 2:665-668.

72. Glassman RH, Sun AY: Biotechnology: identifying advances from the hype. Nat Rev Drug Discov 2004, 3:177-183.

73. Drews J: Strategic trends in the drug industry. Drug Discov Today 2003, 8:411-420.

74. Maeda H, Khatami M: Analyses of repeated failures in cancer therapy for solid tumors: poor tumor-selective drug delivery, low therapeutic efficacy and unsustainable costs. Clin Transl Med 2015, 7:11.

This paper highlights many problems in reductionist drug development derived from high costs. Failed expensively developed cancer drugs may be subject to changes (at times minor) to protocols for example, changes in dosage, route, frequency of drug administration, and the drugs publicised again as ‘new’ approaches to cancer.

75. Prasad S, Gupta SC, Aggarwal BB: Serendipity in cancer drug discovery: rational or coincidence? Trends Pharmacol Sci 2016, 37:435-450.