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Heart-on-a-chip systems: disease modeling and drug screening applications

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Cardiovascular disease (CVD) is the leading cause of death worldwide, casting a substantial economic footprint and burdening the global healthcare system. Historically, pre-clinical CVD modeling and therapeutic screening have been performed using animal models. Unfortunately, animal models oftentimes fail to adequately mimic human physiology, leading to a poor translation of therapeutics from pre-clinical trials to consumers. Even those that make it to market can be removed due to unforeseen side effects. As such, there exists a clinical, technological, and economical need for systems that faithfully capture human (patho)physiology for modeling CVD, assessing cardiotoxicity, and evaluating drug efficacy. Heart-on-a-chip (HoC) systems are a part of the broader organ-on-a-chip paradigm that leverages microfluidics, tissue engineering, microfabrication, electronics, and gene editing to create human-relevant models for studying disease, drug-induced side effects, and therapeutic efficacy. These compact systems can be capable of real-time measurements and on-demand characterization of tissue behavior and could revolutionize the drug development process. In this review, we highlight the key components that comprise a HoC system followed by a review of contemporary reports of their use in disease modeling, drug toxicity and efficacy assessment, and as part of multi-organ-on-a-chip platforms. We also discuss future perspectives and challenges facing the field, including a discussion on the role that standardization is expected to play in accelerating the widespread adoption of these platforms.

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1 Introduction

Cardiovascular disease (CVD) is the leading cause of death worldwide, with an estimated 17.9 million attributable deaths each year,¹ and accounted for 12% of total US health



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expenditures from 2017 to 2018 (\$378 billion US Dollars (USD)).² As such, modeling, preventing, and treating the various subsets of CVD is a highly active area of research.³ While therapeutic intervention has improved patient outcome and quality of life, numerous challenges remain unaddressed. For instance, the hospitalization rate in the US for acute myocardial infarction (AMI), which is caused by an unstable ischemic syndrome,⁴ has declined significantly over the last decade and beyond.² Despite these improvements, post-ischemia reperfusion injury leaves patients especially susceptible to future heart dysfunction/failure,⁵ and, despite promising pre-clinical studies, very few, if any, clinical treatments are available to improve disease prognosis.⁶

The most likely culprit for the poor translation of therapies from pre-clinical studies to widespread clinical usage is the disconnect between the animal models used in pre-clinical studies and human physiology. For example, in the context of the heart, murine models have provided valuable mechanistic insight into heart aging and injury response.⁷ However, the typical heart rate for a mouse is 400 to 650 beats per minute (BPM),⁷ nearly 7 to 10 times that of a human. Furthermore, the predominant isoform of myosin heavy chain (MHC) is α -MHC (encoded by *MYH6*) in fetal humans and β -MHC (encoded by *MYH7*) in adults, although this is reversed in mice.⁸ While these are just a selection of physiological differences between animals and humans, they raise the question regarding the relevance of animal models in pre-clinical assessments. In fact, of the novel drugs that do make it passed pre-clinical animal testing, approximately 89% fail in human clinical trials, oftentimes due to unforeseen toxicity.^{9–11} Even after making it to market, many drugs are removed or recalled due to adverse side effects, with cardiovascular toxicity being the second most common cause behind liver toxicity.¹² Not only do these unanticipated side effects have a tremendous impact on human health and well-being, they come at significant cost to pharmaceutical companies as well. Drug development already takes over 10 years from initial synthesis to completion of clinical trials and costs over \$1 billion USD for a single drug. Recalls can add to this, with a high-profile example being the \$8.5 billion USD spent by a pharmaceutical company for legal settlements after the removal of one of their products from the market.¹⁰

Given the extensive resource requirement for drug development and limited translatability of current animal models, clinicians, pharmaceutical companies, and policymakers are actively searching for a human-relevant pre-clinical alternative to animal models. To address the setbacks of animal models, *in vitro* microphysiological systems, often referred to as “organ-on-a-chip (OoC)”, that have the potential to replicate human physiology and pharmacology have emerged as a potentially viable alternative.¹³ An organ-on-a-chip is, by definition, “a subset of microphysiological systems that replicates one or more aspects of an organ’s *in vivo* dynamics, functionality, structure, and/or (patho)physiological response(s) of multiple cell types integrated within a non-biological

platform”.¹⁴ The synergistic convergence of microfabrication, microfluidics, and tissue engineering has accelerated the development of OoCs in recent times. Microfluidic chips allow for the transport and manipulation of small volumes of liquid, benefiting tremendously from microfabrication processes, rigorously developed in the semiconductor industry. Microfabrication enables the precise spatial arrangement of cell culture chambers, fluidic channels, and analytical components, such as electrodes or cantilevers for real-time monitoring of tissue behavior. To augment these benefits, improvements in tissue engineering have enabled the generation of more complex microtissues comprised of multiple cell types. Furthermore, the advent of human-induced pluripotent stem cells (hiPSCs) has opened up the possibility of tailoring these platforms to individual patients with unique genetic profiles. hiPSCs are especially crucial for the development of heart-on-a-chip (HoC) due to the difficulty of acquiring human heart samples and the limited ability of cardiomyocytes to divide. Overall, these systems have demonstrated the ability to model key aspects of cardiopathologies,^{15,16} tissue regeneration,¹⁷ and pharmaceutical response,¹⁸ oftentimes agreeing well with clinical results.

In this review, we discuss recent reports and advances in HoC systems, engineered cardiac tissue, and cardiac microphysiological systems. We first present a brief overview of cardiac physiology followed by a more in-depth discussion of the major components that make up an organ-on-a-chip. We then focus on two key applications that could benefit from the use of HoCs in addition to or in place of animal testing, namely disease modeling and drug discovery—which includes drug efficacy testing and cardiotoxicity assessment. Finally, a discussion of standardization within the OoC community follows and transitions into the conclusion and future outlook.

2 The heart: a biological pump

The heart is the pump and main organ of the cardiovascular system (Fig. 1). Circulating blood provides tissues and organs with nutrients and oxygen.⁴⁴ The heart is comprised of four chambers which act as reservoirs for blood: the left and right atria in the upper portion of the heart, and the left and right ventricles in the lower portion, separated by the atrial or ventricular septum, respectively.⁴⁵ Oxygen-deficient blood from tissue and other organs enters the right atrium through the *vena cava*.⁴⁵ The oxygen-deficient blood is pumped through the pulmonary artery to the lungs, where it releases waste products (e.g. carbon dioxide) and gets re-oxygenated. The oxygenated blood flows from the lungs back to the heart, where it is circulated to the rest of the body.⁴⁵

To facilitate the circulation of blood throughout the body, the heart wall is comprised of three specialized layers: the endocardium (innermost layer), the myocardium (middle layer), and the epicardium (outermost layer). The endocardium is primarily comprised of endothelial cells, which line the inner walls of the heart’s valves and chambers.⁴⁶ The endocardium is

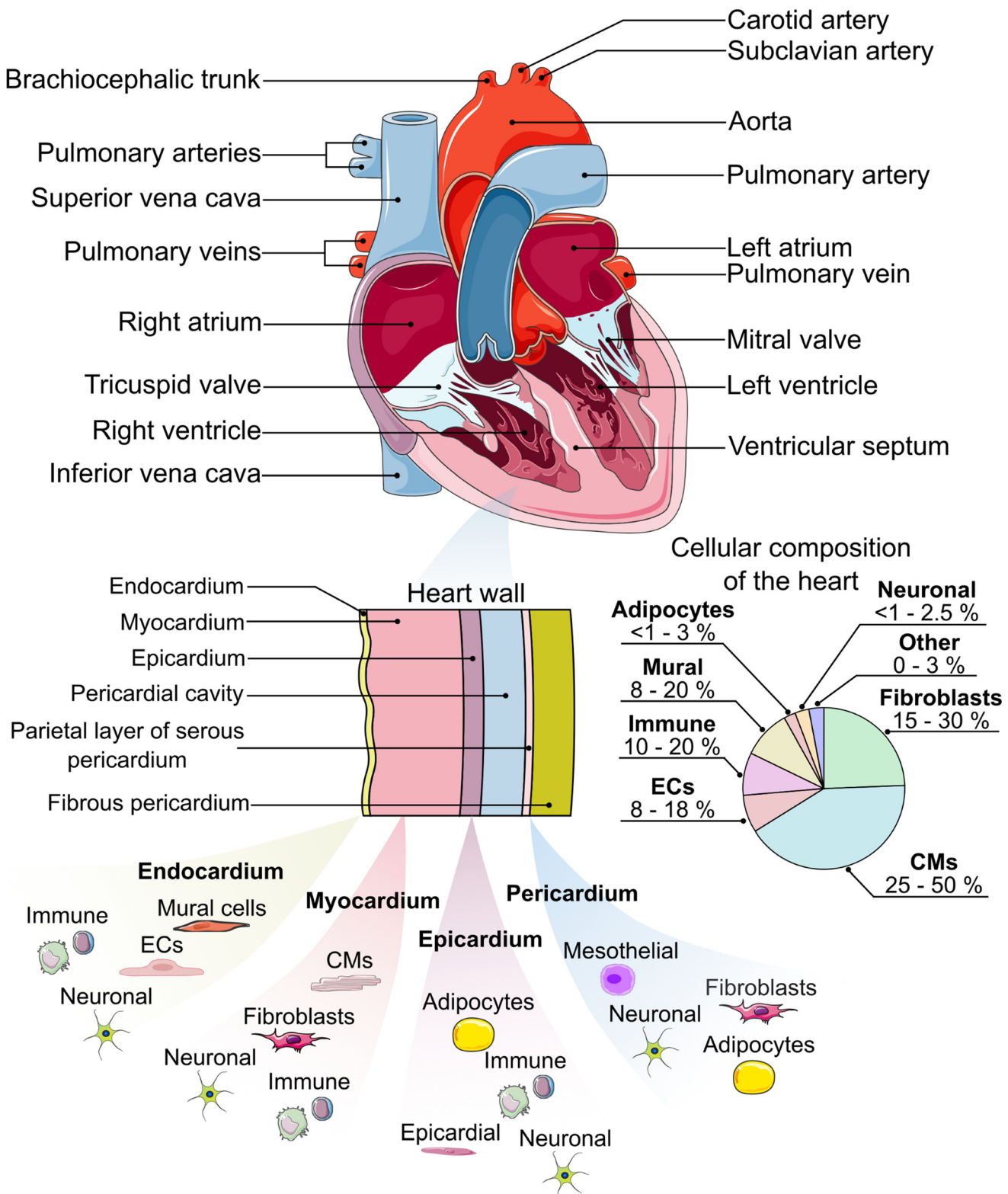


Fig. 1 An anatomical schematic of the heart and heart wall with the main constituent cell types in each layer.¹⁹⁻³⁷ Cellular composition ranges are estimated from the literature.³⁸⁻⁴⁰ ECs = endothelial cells, CMs = cardiomyocytes. Some components of this figure are used under a Creative Commons license.⁴¹⁻⁴³

a thin layer and serves as the barrier between the blood and the remainder of the heart wall. The myocardium is primarily made

up of cardiac fibroblasts, which support the three-dimensional structure of the heart, and cardiomyocytes, which are

responsible for the mechanical contraction of the heart. Cardiomyocytes make up the largest percentage of the heart by volume ($\approx 75\%$), while they are similar to fibroblasts in percentage by number ($\approx 30\text{--}35\%$),³⁹ although other reports have put CM percentages near 50%.³⁸ It is worth noting that the distribution of cell types, especially cardiomyocytes and fibroblasts, varies between heart chambers and between males and females.^{38,40} The epicardium, which includes a layer of mesothelial cells,⁴⁷ plays an important role in paracrine signaling and contributing cardiac progenitor cells during cardiogenesis and post-injury repair.^{48,50} Lastly, the heart is enclosed within the pericardium,^{19,46} which provides lubrication, support for the heart, and protection from blunt force injury and infection.⁵¹

In addition to the major cell types indicated previously, the heart also contains a number of other important cell types that are critical for its function. Specialized cardiomyocytes in the His-Purkinje system, sinoatrial node, and atrioventricular node make up the electrical system of the heart, which is responsible for initiating and transmitting the electrical impulses that enable synchronized contractions.^{46,52,53} Mural cells, including pericytes and vascular smooth muscle cells, help to provide support for the vasculature.⁵⁴ Adipocytes are present in the epicardium and pericardium and assist in protecting the underlying myocardium.⁵⁵ Neuronal cells help to form the cardiac nervous system.⁵⁶ Finally, a variety of immune cells, including macrophages, monocytes, neutrophils, B cells, natural killer (NK) cells, mast cells, and T cells, can also be found in the heart.^{38,39} These cells respond to, for example, infection, myocardial infarction, non-ischemic heart failure, and the presence of toxic compounds.⁵⁷ When designing a HoC model, it is crucial that the appropriate cell type(s) are included to ensure the model captures the native physiology and/or pathology expected in the human body.

The human heart undergoes extensive changes throughout its course of development.⁵⁸⁻⁶⁴ When establishing a HoC model, it is important to consider the maturity of the cells being used. This is especially relevant for cardiomyocytes, which show significant differences in metabolic, structural, and electrical properties between fetal-like and adult cells.^{65,66} Importantly, the proliferative ability of cardiomyocytes declines as cells mature, which can make generating enough cells for *in vitro* experiments a challenge. Structurally, fetal-like cardiomyocytes are typically round and small (5 to 10 μm in diameter) and mono-nucleated, whereas adult cardiomyocytes are larger and elongated (100+ μm long) and most are multi-nucleated.^{65,67-69} The organization and length of sarcomeres also increases as cardiomyocytes mature,⁷⁰ which contributes to an increase in the contractile force generated per cell.⁷⁰⁻⁷² Fetal cardiomyocytes also show differences in myofibrillar isoform expression compared to their more mature counterparts, including titin (N2BA to N2B),^{73,74} troponin (slow skeleton troponin I (ssTnI) to cardiac troponin I (cTnI)),^{75,76} and less α -MHC.⁷⁷ Moreover, the metabolic pathway switches from glycolysis in fetal cardiomyocytes to oxidation of fatty acids in adult cardiomyocytes,⁵⁸ which could impact the response to

drugs and cardiotoxic compounds.⁷⁰ From an electrophysiological standpoint, fetal cardiomyocytes demonstrate a lower membrane potential magnitude (-60 mV vs. -90 mV)⁷⁸ and capacitance,⁷⁹ along with a slower conduction velocity.⁷⁰ Altogether, the degree of cell maturation must be considered when translating results and conclusions from heart-on-a-chip models to human beings.

3 Heart-on-a-chip: components and assembly

HoC systems are comprised of four major sub-systems: the microfluidic architecture (Fig. 2a-c), the cell tissue(s) and associated matrices, environmental controls (e.g. O_2 gradient, drug delivery, mechanical actuation/stimulation) (Fig. 2d-f), and analytical components (e.g. in-line biochemical sensors, electrodes) (Fig. 2g-i).⁸⁰⁻⁸² The conceptual design and construction of a HoC begins with a clear understanding of the target objective(s) and question(s) to be answered,⁸³ which could preclude certain sub-systems. For instance, a key characteristic of a system intended to model myocardial infarction would be a stable and controllable oxygen gradient, whereas a system designed to quantify contractile forces would perhaps emphasize recapitulating the mechanical properties of the cellular microenvironment. As such, the design and engineering of each component within a HoC system is very much interdependent on the others. In this section, each of the four components are discussed in more detail with the goal of providing the reader with a basic understanding of the materials, fabrication techniques, and auxiliary features that are needed when constructing a HoC system.

3.1 Microfluidics

The field of microfluidics deals with the manipulation of small volumes of fluid in microscale (*i.e.* 1–1000 μm in height or width) channels.^{84,85} Microfluidics is applicable across a range of fields, from chemical synthesis to bioanalysis. Microfluidics offers a number of key advantages that make it appealing for organ-on-a-chip systems in particular. Microfluidic systems are inherently dynamic whereas traditional 2D cell culture is largely a static process. The small fluid volume (on the order of nL to μL) minimizes the use of expensive reagents and human samples, and fewer cells are needed for experiments.⁸⁶ The compatibility with many microfabrication methods also allows for the creation of complex chip architectures that can house one or more organ chambers and are capable of cell sorting, fluid mixing, and downstream analysis, all in a device with a small footprint.⁸⁷ Furthermore, microfluidic chips are also capable of mimicking the cellular microenvironment, including physiological shear stress, periodic mechanical strain, and chemical gradients.

While the use of microfluidics in organ-on-a-chip systems has come a long way, one key challenge hindering its widespread adoption is the choice of material(s).⁸⁷ Common materials utilized in the microfluidic structure of HoC systems

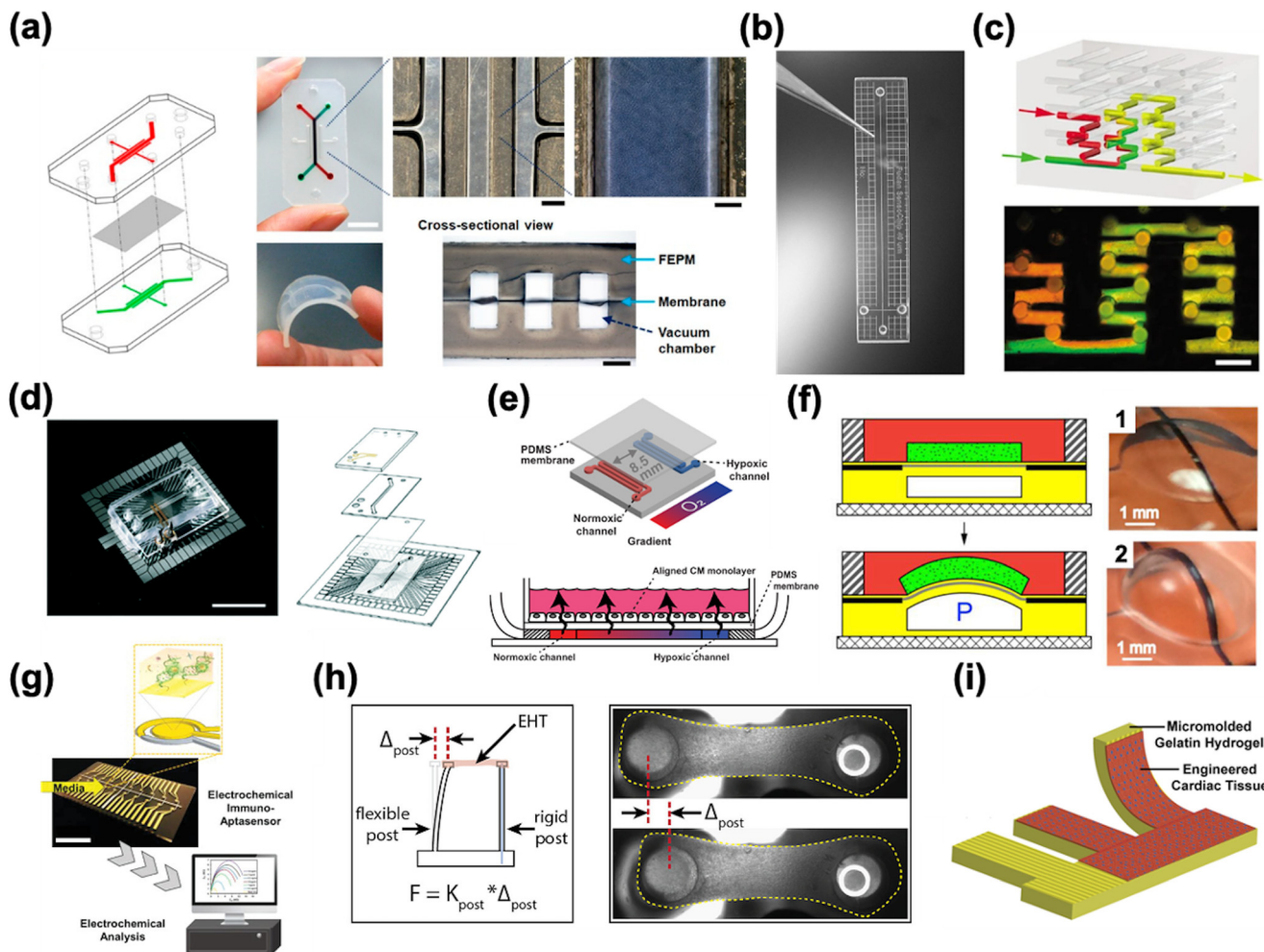


Fig. 2 Exemplary microfluidic chips fabricated using a) a tetrafluoroethylene-propylene elastomer to prevent small molecule absorption (reproduced from ref. 90 under Creative Commons License CC BY 4.0), b) poly(methylmethacrylate) (PMMA) (reproduced from ref. 101 with permission from John Wiley and Sons), and c) a 3D printer (reproduced from ref. 110 with permission from Springer Nature). d) A microfluidic chip combined with a microelectrode array (MEA) (reproduced from ref. 160 with permission from the Royal Society of Chemistry). e) A microfluidic chip with two adjacent channels (one normoxic and one hypoxic) beneath a polydimethylsiloxane (PDMS) membrane for establishment of an oxygen gradient to mimic the infarct border zone (reprinted from ref. 146 under Creative Commons NonCommercial License 4.0 (CC BY-NC)). f) A pressure-actuated membrane with an integrated carbon nanotube (CNT)-based strain sensor for on-chip mechanical conditioning of cardiac cells (1 = resting, 2 = actuated) (reproduced from ref. 161 with permission from Elsevier). g) An electrochemical immuno-aptasensor array for biomarker quantification (reproduced from ref. 162 with permission from John Wiley and Sons). h) A two-pillar construct for monitoring the mechanical behavior of cardiac tissue based on the deflection of the flexible post (reproduced from ref. 163 with permission from Elsevier). i) An example of a muscular thin film array (MTF), which quantifies mechanical behavior of cardiac tissue by measuring the curvature of the flexible cantilever as the tissue contracts and relaxes (reproduced from ref. 164 with permission from Elsevier).

include polydimethylsiloxane (PDMS), poly(methyl methacrylate) (PMMA), polyurethane (PU), polystyrene (PS), and other elastomers.^{88–91} Pattern transfer to these materials can be done using a variety of techniques,⁹² including soft lithography,⁹³ photolithography,⁹⁴ hot embossing,^{95,96} and 3D printing,^{97,98} to name a selection. PDMS is widely used for prototyping and lab-scale experiments due to the ease with which it can be cast and molded. Furthermore, its optical transparency and gas permeability make it well-suited for prolonged cell culture and monitoring.⁸⁸ PDMS chips can also be integrated with auxiliary components, such as porous membranes for cell migration studies for example, *via* a plethora of bonding techniques. Nonetheless, PDMS is

relatively expensive and readily absorbs many small molecules, making applications such as drug discovery particularly challenging.¹³ As a result, other materials are being explored as alternatives to PDMS. PMMA, PS, and polycarbonate (PC) are all biocompatible, inexpensive, ubiquitous materials used in cell culture system and microfluidics.^{99–103} The challenge with these materials is they are relatively difficult to microfabricate in an academic lab, autofluoresce, and can be difficult to interface with peripheral equipment, such as pumps.^{104–106} PU is a less-explored material that addresses many of the drawbacks associated with PDMS, all while maintaining the advantageous properties, including optical transparency for microscopy.^{107,108}

It is worth briefly highlighting recent efforts to 3D print microfluidic components, which is widely seen as a new paradigm in manufacturing organ-on-a-chip systems.¹⁰⁹ 3D printing is an additive process that involves the layer-by-layer deposition of a material in a precisely programmed fashion. 3D printing is commonly used to quickly prototype and fabricate molds for casting PDMS, which are otherwise prepared using traditional microfabrication processes, such as photolithography. However, due to the limited scalability of PDMS, researchers are exploring the direct printing of microfluidic components rather than printing of soft lithography templates. Direct 3D printing of microfluidic components enables truly three-dimensional, multi-planar fluidic constructs (e.g. a helix),¹¹⁰ rather than two-dimensional patterns that are projected into the third dimension. Furthermore, 3D printing is largely an automated process that involves significantly less manual labor when assembling organ-on-a-chip devices. Despite the advantages 3D printing offers, critical technological challenges remain, including the resolution limit of the printer, the cytocompatibility of the resin materials, low throughput, low gas permeability, and ensuring water-tight sealing of the channels. Nonetheless, 3D printing offers tremendous prospect for bringing organ-on-a-chip products to market.^{84,111}

3.2 Cell sources and matrices

Before the HoC system can be engineered and constructed, careful consideration of the type(s) of cell(s) to be incorporated is necessary. Animal-derived cardiac cells, typically from the hearts of rats or pigs, are one potential source of cells. The handling, isolation, and cultivation of animal-derived cells is well-established and reproducible. While animal models and animal-derived cells have provided insight into heart development, they can be less effective at modeling cardiac diseases and adverse drug effects due to differences between human and animal physiology and pharmacodynamics.^{112,113} The advent of human embryonic stem cells (hESCs) and, more recently, human-induced pluripotent stem cells (hiPSCs) has enabled *in vitro* heart models with cells of human origin.^{112,114} hiPSCs, in particular, alleviate many of the ethical and legal issues surrounding the use of hESCs,¹¹⁴ and therefore, it is worth emphasizing the advantages hiPSCs offer. The isolation of human cardiac samples is difficult, if not impossible, and the collected cardiomyocytes do not easily divide, making culture maintenance and expansion challenging. On the contrary, hiPSCs can be derived from somatic cells obtained through trivial means, such as a skin or blood sample. hiPSCs readily proliferate, making them amenable to conventional cell culture practices. Additionally, hiPSCs possess the genetic profile of their source, opening up the potential for patient-specific drug screening and personalized disease modeling.

hiPSCs are oftentimes differentiated into cardiomyocytes (hiPSC-CM), which is now a well-established procedure, given the large volume of the heart they comprise ($\approx 75\%$ by volume, $\approx 30\%$ by number).^{115,116} The use of hiPSC-CMs has enabled the evaluation of various cardiotoxic compounds and

pathological environments on contraction dynamics *in vitro*. To create a more holistic *in vitro* model with multiple cell types, methods for differentiating hiPSCs into other cell types, such as endothelial cells (EC) and cardiac fibroblasts (CF), which are responsible for producing the extracellular matrix (ECM) *in vivo*, are being developed.¹¹⁷⁻¹¹⁹ Strategically incorporating different cell types to more accurately represent the heart *in vivo* could enable more comprehensive studies of cell-cell communication within the heart. However, the controllable placement and selective differentiation of multiple cell types for *in vitro* modeling has proven to be challenging.^{120,121} Perhaps the most pressing challenge facing the widespread adoption of hiPSC-based *in vitro* models is the immaturity of hiPSC-derived cardiac cells.^{8,59,122,123} It is known that mature CMs, for instance, show distinctly different mechanical, morphological, and electrophysiological characteristics compared to immature CMs. Such differences can directly impact how the cells respond to certain stimuli and stressors, leading to an incomplete understanding of the model.⁸ Guided by the mechanisms of cardiac development *in vivo*, methods for cardiac maturation in HoC systems are being explored. Perhaps the most simple method for maturation is to culture the CMs for a long period of time (>6 months), which leads to cells that more closely resemble the phenotype and of mature CMs.⁸ While easily implemented, such long culture duration is not practical for the applications targeted by HoC systems. Other maturation methods include co-culturing with other cardiac cells, patterned cell seeding, biochemical means, and mechanical or electrical stimulation/conditioning, which are discussed further in section 4.1.1.

Depending on the type(s) of cell(s) to be incorporated in the HoC system, careful selection of the ECM is paramount.¹²⁴⁻¹²⁶ For example, seeding undifferentiated hiPSCs or hESCs requires an ECM that can facilitate the subsequent differentiation. Likewise, seeding hiPSC-CMs requires an ECM suitable for tissue formation. Most commonly, ECMs incorporate proteins, hydrogels, polycations and polyanions, growth factors, and various small molecules.¹²⁷⁻¹³⁰ Common proteins incorporated in ECMs include various isoforms of laminin, collagen, fibronectin, and vitronectin.^{129,131} Matrigel,¹³² a common commercially available ECM from Corning that is comprised of 60% laminin-111 and 30% collagen-IV, is often used for hESC and hiPSC culture. However, because Matrigel is derived from animals (Engelbreth-Holm-Swarm mouse sarcoma cells), it lacks chemical definition, which can lead to varying concentration of proteins and a lack of reproducibility between studies.¹³³ These discrepancies have motivated investigations into whether particular protein isoforms can support hESC and hiPSC growth and differentiation. Laminin-111, -511, and -521 have had the most success in supporting the growth and efficient differentiation of pluripotent hiPSCs and hESCs, especially laminin-521 combined with laminin-211 or -221.^{115,134,135} However, on their own, laminin-211 and -221 were unable to support pluripotent stem cell growth and differentiation.¹³⁵ Collagen I has also been shown to be a poor ECM for pluripotent stem cell differentiation into CMs.¹³⁴

Fibronectin and vitronectin are both glycoproteins expressed within the ECM of the heart. Nonetheless, hiPSCs have shown poor growth on fibronectin,¹¹⁵ making subsequent differentiation into cardiomyocytes a challenge. hiPSC-CMs reseeded on fibronectin coated substrates were able to attach but showed inferior contractile behavior compared to those seeded on laminin.¹³⁶ While vitronectin has been shown to support hiPSC differentiation to CMs,¹³⁷ it does not appear to be the optimal ECM material. CM yield was lower on vitronectin when compared to Matrigel and laminin.¹¹⁵ Moreover, Sung *et al.* reported a slower beating frequency, lower cTnT expression, and shorter sarcomere length for hESCs differentiated on vitronectin compared to Matrigel and other common ECM materials.¹³⁴

3.3 Environmental controls

The purpose of introducing environmental control into the HoC system is to better recapitulate the conditions experienced by the cells *in vivo*. HoC models can be studied using a standard cell culture incubator, with a temperature of 37 °C, atmospheric oxygen level of 21%, and high relative humidity (>90%). The bicarbonate buffer of many common culture media requires a CO₂ concentration of 5% to maintain a physiologically relevant pH value at or near 7.4.⁹¹ If the HoC system will be maintained outside of a standard culture incubator, the media can be preconditioned in an incubator before adding to the chip to ensure appropriate pH. HEPES buffer may also be added to maintain physiologically relevant pH values.^{91,138} Inside the chip, environmental controls can be used to promote cardiomyocyte maturation, dictate cell structure, location, and orientation, and replicate pathological conditions. Environmental controls can be either static or dynamic and include electrical/mechanical/biochemical stimulation, spatial and temporal chemical gradients, and templated culture regions, among others.

Electrical stimulation, or pacemaking, can be incorporated into a HoC system through the use of microfabricated electrodes.^{139,140} Applying a periodic voltage pulse to the electrodes can regulate cardiomyocyte contractions. Mechanical stimulation, on the other hand, can be accomplished through a variety of methods.¹⁴¹ Typically, it is accomplished by growing cardiomyocytes on a flexible material that can be actuated periodically, which mimics the mechanical contractions experienced *in vivo*.¹⁴² To realize this actuation, specific vacuum channels are oftentimes incorporated within the HoC, causing the flexible growth substrate to bend when vacuum is applied, and flatten when vacuum is relaxed. Biochemical stimulation involves introducing a stimulant into the culture chamber to increase or decrease beating rate, such as isoproterenol.^{143–145}

One key advantage of incorporating microfluidics into HoC systems is having control over the spatial distribution of (bio)chemicals of interest. Chemical gradients are often encountered in the body, for example, in the context of myocardial infarction, where a portion of the heart is deprived of the necessary oxygen, and pharmacokinetics and pharmacodynamics, which are important for drug efficacy

evaluation. These gradients can be established through the use of two or more independent channels with differing levels of the chemical of interest.^{146,147} Porous and/or permeable membranes may also be incorporated into the HoC system to help establish the necessary gradients and model chemical transport through a barrier.^{146,148} For instance, PET membranes can be sandwiched between two PDMS chips to allow for cross-talk between upper and lower channels.¹⁴⁹ PDMS can also serve as a membrane to control oxygen levels given its excellent gas permeability. Incorporating polymer membranes requires the surface to be functionalized to promote bonding with the microfluidic layer. Plasma treatment and arc discharge are useful for generating reactive oxygen groups on the surface of both the polymer membrane and microfluidic layer. Silane chemistry is another common functionalization strategy that can be used to introduce irreversible amine-epoxy or O–Si–O bonds between the membrane and microfluidic layer.^{150–152}

Cell patterning is a technique for controlling the spatial position and orientation of cells on a substrate.¹⁵³ This technique often involves using topographical cues to promote a particular cell alignment or morphology,⁶⁰ which is especially relevant when culturing cardiomyocytes to promote tissue anisotropy and uniaxial contraction as seen *in vivo*.^{154–157} Orienting cells can be accomplished by patterning the extracellular matrix in such a manner that promotes cell growth along a particular direction.¹⁴⁶ Engineering a cell “template” through the use of physical barriers, guides, and pillars can also be used to direct cell growth and orientation.^{139,158,159} Ensuring biologically relevant orientation of the tissue with a HoC helps to better replicate the *in vivo* mechanical properties and subsequent response to stimuli and stressors.

3.4 Analytical components

One of the major advantages of organ-on-a-chip systems is the ability to integrate in-line analytical tools within the chip. Generally, researchers are interested in characterizing electrophysiology, contractile force, secreted biomarker dynamics, or mechanical properties of the cells/tissue.¹⁶⁵ Examples of in-line analytical tools include microelectrodes, electrochemical biosensors, field-effect transistor (FET) biosensors, elastic cantilevers, optical microscopy, optical sensors (e.g. surface plasmon resonance (SPR)), magnetic sensors, and piezoelectric biosensors, which are chosen based on the desired properties to be measured.^{166,167} Some examples have the added benefit of acting as both an analytical component and environmental control. For instance, a microelectrode could be used to electrically stimulate the tissue, thereby expediting maturation, while also allowing for electrophysiological measurements to be made in tandem.

The preparation and integration of an analytical component begins by determining the characteristic(s) to be measured. Microelectrodes, including electrical and FET biosensors, are often used for probing the electrophysiology of entire tissue down to individual cells by recording changes in impedance,

potential, *etc.*^{168–173} These measurements require cells to be grown or seeded directly in contact with the electrode. Planar microelectrodes and microelectrode arrays are typically prepared *via* semiconductor fabrication methods, enabling highly controllable electrode size, geometry, and density. The most common substrate for microelectrode fabrication is glass given its low cost, biocompatibility, optical transparency, and ease with which it bonds to PDMS. Probing individual intracellular electrophysiology generally requires nano- or micro-pillars that can penetrate the cell membrane without damage.¹⁷⁴ Again, semiconductor fabrication methods allow for precise control over pillar location and size. The pillars are made from an inert metal (*e.g.* Pt) to ensure adequate conductivity and minimal chance of cytotoxicity. By fabricating arrays of microelectrodes or micropillars, tissue properties can be sampled at multiple locations and time points, which improves data robustness and allows for real- or near real-time measurements. Moreover, these arrays can be easily integrated with peripheral data processing and readout equipment as well. Electrochemical and FET biosensors can also be used to for biomarker quantification. To ensure sufficient selectivity towards the target biomarker, the sensor surface is functionalized with a bioreceptor, such as an antibody, aptamer, nucleic acid, *etc.* Common electrode materials include metals, conductive polymers, and nanomaterials. These sensors are typically placed downstream from the cell culture chamber to capture secreted biomarkers, such as proteins and small molecules.^{162,166,175}

Mechanical properties, such as contractile force, can be probed using two-pillar constructs,^{176–178} flexible cantilevers,^{164,179,180} and piezoelectric materials.¹⁸¹ The two-pillar structure typically serves more as a guide for tissue formation, rather than a sensor *per se*, and it often requires a microscope to visualize contraction dynamics. Likewise, the flexible cantilever often requires optical visualization to quantify cantilever displacement, which is then used to extract mechanical information. However, some recent examples have integrated electrical strain sensors with the flexible cantilever to monitor contraction mechanics.^{180,182,183} Similarly, permanent magnets can be incorporated into one of the two pillars to generate an electromagnetic signal to characterize mechanical contractions.¹⁸⁴ Piezoelectric materials inherently convert mechanical energy into an electrical output, making these materials well-suited for recording contraction profiles from cardiomyocytes.¹⁸¹

4 Applications of heart-on-a-chip technologies

Once the HoC system has been assembled, the target cell type(s) are cultured in the region of interest. The cells are allowed to grow for the desired duration to ensure the appropriate degree of maturation. Perhaps the three most common and apposite applications for HoC devices are disease modeling, cardiogenesis, and drug/therapeutic screening. The particular application and physiology will dictate the complexity of the system, which in the simplest case contains one cell type. More advanced models could require two or more cell types, for

example to study the cardiotoxicity of chemotherapeutic drug metabolism in another organ. In the following section, examples of HoC systems for disease modeling and drug screening are explored in more detail.

4.1 Disease modeling and cardiac maturation

Control over the cellular microenvironment and spatiotemporal (bio)chemical distribution makes HoC platforms well-suited for *in vitro* disease modeling and studying cardiac maturation. Pathological conditions can be modulated in a precise manner while on-board analytical tools enable real-time measurements of tissue functionality, which is difficult to achieve with endpoint measurements commonly used in static 2D cultures. Furthermore, the advent of hiPSC-derived cardiomyocytes has provided an opportunity to model diseases with cells containing an individual's unique genetic profile, thereby providing a more personalized understanding of disease progression. To that end, HoC systems have been leveraged to study cardiac maturation as well as model a range of cardiomyopathies, myocardial infarction, ischemia/reperfusion injury (IRI), and fibrosis, discussed in more detail in the following section.

4.1.1 Cardiomyocyte maturation. The importance of hiPSC-derived cardiac cells in HoC platforms cannot be overstated. While animal models and animal-derived cell lines have played a crucial role in many of these applications, they do not adequately recapitulate the structural and physiological properties of the human heart.¹⁴¹ The ability to easily obtain hiPSCs and readily generate a large number of patient-specific cardiac cells, including cardiomyocytes, fibroblasts, and endothelial cells, has been a game changer for *in vitro* applications. However, despite these advances, hiPSC-derived cells oftentimes display an immature/fetal-like phenotype that fails to fully mimic the human adult heart.^{59,185} For instance, hiPSC-CMs typically rely on glycolysis for energy production, which resembles embryonic CMs, rather than lipid oxidation commonly seen in adult CMs.¹⁸⁶ To improve the relevance and translatability of *in vitro* models, various techniques for promoting maturation, including mechanical, electrical, (bio)chemical, and cell co-culture, have been investigated.^{8,123,141}

4.1.1.1 Long-term culture. Perhaps the most straightforward method to promote CM maturation is long-term culturing. Lundy *et al.* cultured hiPSC- and hESC-derived CMs for over 100 days and noted significant increases in sarcomere length, myofibril organization, cell size, and percentage of multi-nuclear cells.⁷² Subsequently, the calcium handling and contractile function were improved, explained in part by a shift from the α - to β -isoform of myosin heavy chain as evidenced by changes in gene expression. Similar morphological shifts were seen by Snir *et al.* with hESC-CMs after 36 days in culture.⁶⁸ This observation aligns with genomic data reported by Piccini *et al.* that showed stark differences in the expression of structural markers, such as *MYH6*, *MYL7*, and *MYL4*, between 1 week and 8 week old hPSC-CMs.¹⁸⁷ The differences between 4 week and 8 week

old hPSC-CMs, however, was minimal, indicating much of the maturation takes place in the initial 4 weeks of culture.

4.1.1.2 Biochemical stimulation. While long-term culture is a seemingly straightforward means to promote CM maturation, the long time requirements may not be suitable in all settings. Biochemical cues appear to be a suitable alternative method to promote CM maturation. Tailoring the level of fatty acids, glucose, and galactose in the culture media can have substantial impacts on CM maturation.^{188–192} This promotion of maturity is due to a shift in the energy metabolic pathway from glycolytic to oxidative, which more closely captures that seen in the adult heart. However, media supplemented entirely with fatty acids and no carbon source appeared cytotoxic.¹⁹⁰ This effect could be reversed upon incorporation of galactose to improve the oxidative capacity of the cells, although others have reported galactose incorporation alone was not sufficient for improving cell viability.¹⁸⁸ Hormones have also demonstrated the ability to promote cardiac maturation. Yang *et al.* investigated the effect of tri-iodo-l-thyronine (T3), a thyroid hormone, on cardiac maturation.¹⁹³ The incorporation of T3 into the culture medium promoted physiological hypertrophy, contractile force, cell size and anisotropy, and increased mitochondrial respiratory function, all of which are signs of a more mature state. Parikh *et al.* also investigated T3 combined with the glucocorticoid, dexamethasone, on cardiac maturation, which increased the density of T-tubules and improved Ca^{2+} synchronization.¹⁹⁴ Supplementing culture media with growth factors has been implicated as a way to improve maturity. Rupert and Coulombe studied two growth factors crucial to cardiac development, insulin-like growth factor 1 (IGF-1) and neuregulin-1 β (NRG1).¹⁹⁵ IGF-1 was found to drastically increase CM proliferation on its own, while NRG1 promoted a more mature metabolic phenotype. However, timing appears to be crucial given that simultaneous administration of both growth factors negated some of the positive effects, such as improved CM proliferation.

4.1.1.3 Mechanical conditioning. Mechanical conditioning has been reported as a means to promote cardiac maturation.^{197–203} The goal is to mimic the various forces that cardiac cells and tissue experience during development.^{124,141,204} These include shear stress from blood flow, strain arising from cyclic contraction, stretching as a result of blood pressure changes, and forces caused by changes to the elastic modulus of the tissue. Passive mechanical stimulation was demonstrated by Rao *et al.* who developed a culture substrate with parallel PDMS microscopic grooves coated with fibronectin.²⁰⁵ The grooves helped guide the alignment of hPSC-CMs as seen by more organized sarcomere assembly. Subsequent analysis of Ca^{2+} dynamics generally showed more rapid kinetics for the cells grown on structured substrates as opposed to non-structured. To study the effect of shear force on cardiac maturation, Cruz-Moreira *et al.* created a HoC platform with peristaltic pump capabilities.²⁰⁶ CMs subjected to the highest flow rate (48 $\mu\text{L min}^{-1}$) were structurally and functionally more mature,

demonstrating an upregulation of many genes involved in structure (e.g. *ACTC1* and *MYH6*), intercellular communication (e.g. *GJA1* (encodes connexin 43)), calcium cycling (e.g. *SERCA2*) and metabolism (e.g. *PGC-1A*) and a downregulation of hypoxia-inducible factor 1 alpha (*HIF-1 α*) which suggests improved oxygen and nutrient delivery at higher flow rates. The effect of afterload, which is the systolic load experienced during a contraction, was studied using a two-post system with tunable stiffness (Fig. 3a).¹⁶³ Low to moderate levels (0.1 to 1 $\mu\text{N } \mu\text{m}^{-1}$) of afterload were found to promote cardiac maturation, as indicated by improved contractile function, improvements in Ca^{2+} handling, and genetic markers of maturation.

4.1.1.4 Electrical pacing. Electrical pacing attempts to replicate the constant electrical impulses experienced by CMs *in vivo*.¹²² These impulses are typically applied at frequencies in the range of 0.5 Hz to 5 Hz. Tissue subjected to electrical stimulation has shown evidence of hypertrophy, improved contractile function and calcium handling, a more organized distribution of sarcomeres, and upregulation of genes associated with maturation (Fig. 3b).^{139,140,154,196,207–211} Ronaldson-Bouchard *et al.* compared hiPSC-CMs harvested immediately after first spontaneous contractions (early-stage) and those harvested after 28 days in culture.²¹¹ Each cohort of cells was subjected to either no electrical stimulation, constant stimulation at 2 Hz, or gradual ramping from 2 Hz to 6 Hz at 0.33 Hz per day. Constant stimulation did have some effect on the early-stage cells in terms of gene expression, the gradual frequency ramping of early-stage tissue showed an upregulation of many genetic markers associated with maturation, including *MYH7*, *GJA1*, *NPPA*, and *MAPK1*. Additionally, increases to cell area, sarcomere organization, area of mitochondria, and contractile force were reported as well. However, these changes were not nearly as apparent in late-stage tissue subjected to similar conditioning.

4.1.1.5 Cellular co-culture. Given that CMs comprise only 30% by number of total cells in the heart, co-culture with other cell types, such as fibroblasts, neural cells, endothelial cells, and immune cells has been shown to promote maturation through intercellular interaction and paracrine signaling.^{8,212–226} Co-culture of hESC-CMs with human cardiac fibroblasts (at an optimized 4:1 ratio) in a microfluidic chip containing an array of microposts was found to improve tissue maturity and anisotropy after just 2 weeks (Fig. 3c).¹⁵⁹ Closer inspection revealed that the fibroblasts were capable of proliferating into the surrounding region, perhaps due to the mechanical cues arising from the microposts. Numerous genes associated with maturation were also upregulated for cardiac tissue culture with microposts, including conduction genes (e.g. *HCN1*, *HCN2*), calcium handling genes (e.g. *CAV2.1*, *CAV3.1*, *PLN*), and structural genes (e.g. *GJA5*, *TNNT2*). Co-culture of CMs with endothelial cells has also been shown to improve expression of cardiac maturation markers, including *Cx43*, *TNNI3*, *Kir2.1*, and *CD36*, with a 3:1 CM:EC ratio being most favorable.²²⁶ Moreover, this ratio showed improved electrophysiological characteristics and sarcomere

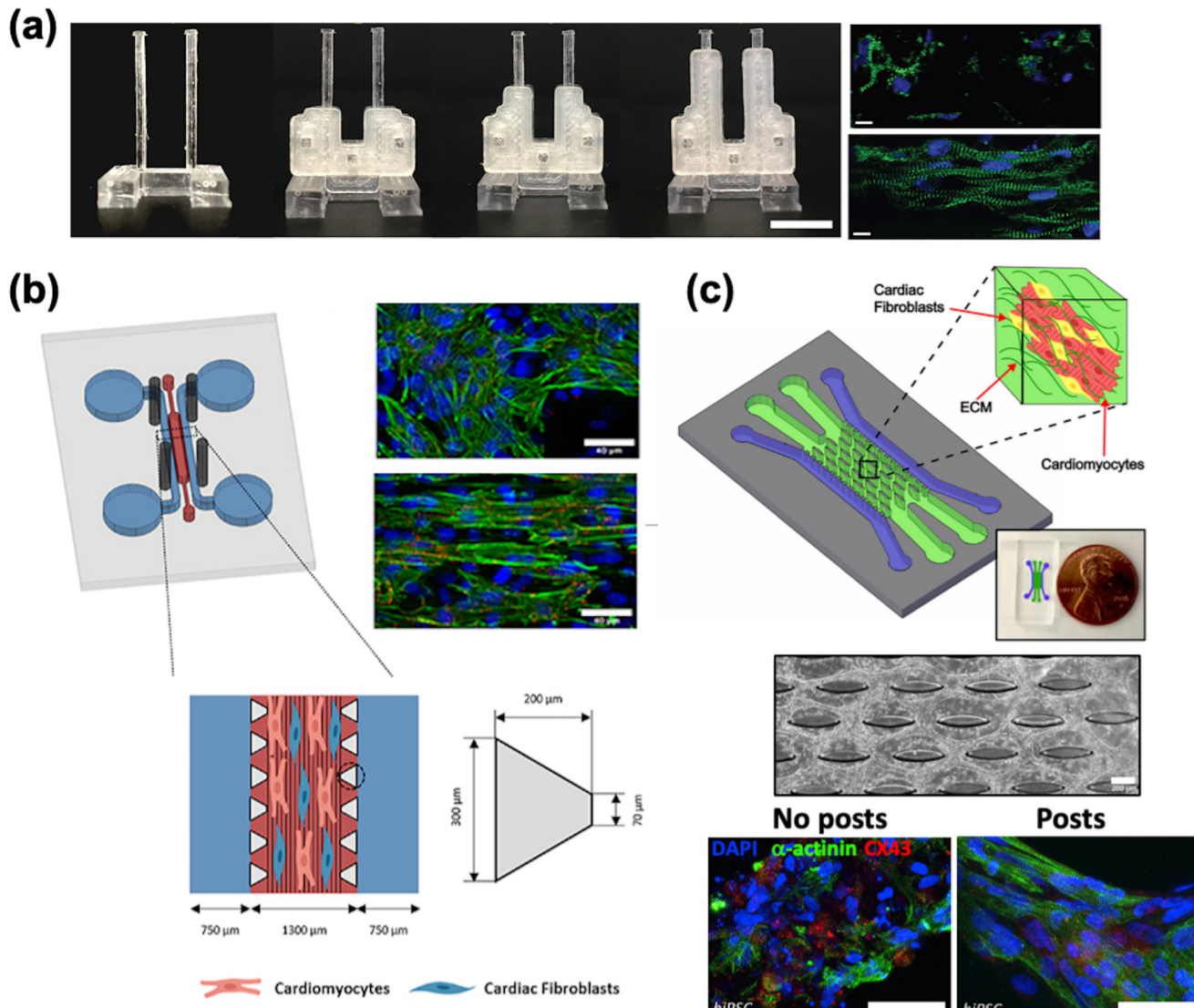


Fig. 3 Cardiac maturation methods. a) A two-pillar construct with braces of varying length to modify the stiffness of the post and resulting afterload. The fluorescence images show more aligned cells grown with afterload (bottom) compared to without afterload (top) (scale bar = 10 μm) (reproduced from ref. 163 with permission from Elsevier). b) A multi-channel PDMS chip with steel electrodes to generate an electric field perpendicular to the channel, leading to improved tissue alignment (reproduced from ref. 196 with permission from IOP Publishing). c) A multi-channel chip with an elliptical micropost array to provide support for tissue growth (scale bar = 200 μm). Immunostaining of cardiac markers shows improved tissue alignment (scale bar = 50 μm) (reproduced from ref. 159 with permission from Elsevier).

organization. While 4 genetically distinct subtypes of CM were identified when hiPSC-CMs were cultured alone, hiPSC-CMs cultured with ECs showed an additional subtype that upregulated genes associated with metabolism and cardiac muscle contraction. Furthermore, the authors examined ligand-receptor complexes which unveiled many intercellular signaling pathways between CMs and ECs, including endothelin-1, ephrin, and vascular endothelial growth factor (VEGF), that are responsible for promoting CM organization, maturation of the vasculature, and the development of the conduction system. Taken together, these results suggest robust intercellular communication between CMs and other cell types and present a means to improve cardiac maturation and biological relevance.

4.1.2 Myocardial infarction and ischemia-reperfusion injury. Myocardial infarction is caused by an occlusion in the coronary artery that results in the oxygen-deprivation of downstream cardiac tissue. Within 20 minutes to 40 minutes, sarcolemmal disruption can occur, followed by mitochondrial changes. Widespread cell death and necrosis occur within hours, even in as little as 20 minutes. Similar phenomena were demonstrated in H9c2 cells using a HoC platform designed by Ren *et al.*¹⁴⁷ Here, the authors developed a system with 4 parallel culture chambers, each capable of sustaining an oxygen gradient across a central channel through the use of carbonyl cyanide-*p*-trifluoromethoxyphenylhydrazone (FCCP)-infused culture media. The presence of an oxygen gradient mimics the “border-zone” established between infarct and normal tissue

in vivo, which has been implicated as a source of arrhythmia. The authors noted alterations to the mitochondrial membrane potential, along with a decrease in average cell size and an increase in the activation of caspase-3 (an indicator of apoptosis), were confirmed using fluorescence microscopy and found to be most drastic at the hypoxic boundary of the channel and largely unchanged at the normoxic boundary. Rexus-Hall *et al.* also developed a HoC device to study the border zone resulting from myocardial infarction.¹⁴⁶ Using an elastic muscular thin film assay (MTF), the authors noted a dysfunction in contraction mechanics, namely a decrease in peak systolic and diastolic stress at the normoxic boundary compared to a uniformly normoxic environment, in agreement with previous reports.^{227,228} Furthermore, changes in the transcriptional profile and a decrease in Ca^{2+} wavefront propagation velocity were found as well, in agreement with previous reports.²²⁹ Similarly, Liu *et al.* reported changes in the contraction dynamics, wavefront propagation, and action potential of HL-1 cells under hypoxic conditions.¹⁷⁴ While their system did not establish an oxygen gradient, it did incorporate both extra- and intracellular electrical sensors. The authors recorded an initial period of tachycardia after removing oxygen from the medium, followed by a decrease in the beating rate to below normoxic levels and the development of an arrhythmic beating pattern.

Restoration of blood flow and oxygen supply after the ischemic episode results in the generation of reactive oxygen species, rapid changes in intracellular pH, and an overload of Ca^{2+} , which can result in further cell death.²³⁰⁻²³⁷ Additionally, due to the poor regenerative ability of the myocardium, post-ischemia tissue reparation results in increased scarring and, subsequently, increased stiffening of the myocardium. However, pre-clinical animal trials have failed to successfully recapitulate the human physiology and, as a result, no clinical treatment for IRI exists despite its ubiquity. To help address the lack of clinically relevant treatments, Chen and Vunjak-Novakovic developed an on-chip IRI model which utilized flexible pillars for hiPSC-CMs to attach and align to.²³⁸ The authors explored four therapeutic strategies using the developed system: ischemic preconditioning, normalization of intracellular pH, minimization of mitochondrial permeability transition pore (MPTP) opening, and reduction of oxidative stress levels. Ischemic preconditioning, whereby the tissue is exposed to brief cycle(s) of controlled ischemia and reperfusion before the primary ischemic or reperfusion episode,^{5,239} was found to improve cell viability after reperfusion, possibly due to the activation of pro-survival kinases during preconditioning.^{240,241} Intracellular pH normalization and minimization of MPTP opening were also found to improve outcomes after reperfusion to varying degrees. On the contrary, treatment with an antioxidant during reperfusion to mitigate oxidative stress levels did not lead to an improved outcome. Clinical trials studying the effect of antioxidants on patients undergoing MI or coronary artery bypass surgery have shown varying results,²⁴²⁻²⁴⁷ which further emphasizes the need for robust pre-clinical *in vitro* models. While most studies focus on the effect of MI and IRI

on cardiomyocytes given their role in the degradation of contractile force *in vivo*, they comprise only 30% of the cell population in the heart. Endothelial cells, fibroblasts, and immune cells make up a substantial portion of the cell population within the heart, and their incorporation in an *in vitro* model can improve physiological relevance. Fibroblasts, for instance, have demonstrated cardio-protective behavior through intercellular signaling.²⁴⁸⁻²⁵⁰ Endothelial²⁵¹⁻²⁵³ and immune²⁵⁴⁻²⁵⁶ cells have also demonstrated cardio-protective and cardio-reparative capabilities through nitric oxide generation *via* endothelial nitric oxide synthase (eNOS) and other anti-inflammatory behavior. In contrast, others have reported pro-inflammatory effects from fibroblasts²⁵⁷ and vascular damage from endothelial and immune cells.²⁵⁸⁻²⁶⁰ Furthermore, the heterogeneity in spatiotemporal and phenotypic distribution of immune cells is extremely complex during and after ischemia but largely dictates the degree of inflammatory and reparative response.^{255,261,262} While it is impossible to completely replicate the complex orchestrated response to MI in a HoC system, the addition of other cell types may lead to more physiologically relevant results and eventually improved clinical outcomes.

To this end, Veldhuizen *et al.* developed a HoC platform with embedded microposts to improve tissue formation using hiPSC-CMs and CFs as model cell lines.²⁶³ The authors examined three environments, namely hypoxic (1% O_2), physioxic (5% O_2), and hyperoxic (21% O_2), and demonstrated some of the hallmarks of an ischemic episode, including altered expression of the key hypoxia-responsive genes, smooth muscle alpha (α)-2 actin (*ACTA2*) and vascular endothelial growth factor A (*VEGFA*). However, no changes in transforming growth factor beta-1 (*TGFB1*) gene expression were found. It is speculated that the addition of macrophages into the platform could change this given they release the proteins $\text{TGF}\beta$ -1 and angiotensin II (ANGII). Signs of contractile dysfunction, reduced cell viability after reperfusion, and increased markers of fibrosis, including α -smooth muscle actin (αSMA) and F-actin were also reported. Interestingly, (αSMA) and F-actin levels did not change when hiPSC-CMs and CFs were co-cultured in 2D on a coverslip, further highlighting the importance of 3D culture models to more accurately replicate human physiology.

Yadid *et al.* also noted a reduction in cell viability and contractile function in human stem cell-derived CMs subjected to hypoxia for 3 h.²⁶⁴ Here, the authors set out to better elucidate the role of endothelial cell-derived extracellular vesicles (EVs) in mitigating the deleterious effects of IRI. EVs are small (typically <300 nm) cell-secreted particles that play an important role in intercellular communication by transporting proteins, lipids, microRNAs, and other genetic material between cells.^{265,266} While the content of an EV is highly dependent on the source and receptor cell type, in general, EVs appear to be cardio-protective or reparative.²⁶⁷⁻²⁷⁷ Yadid and co-workers found that treatment with EVs (ECs are not in co-culture) after IR led to improved viability and contractility with the treated

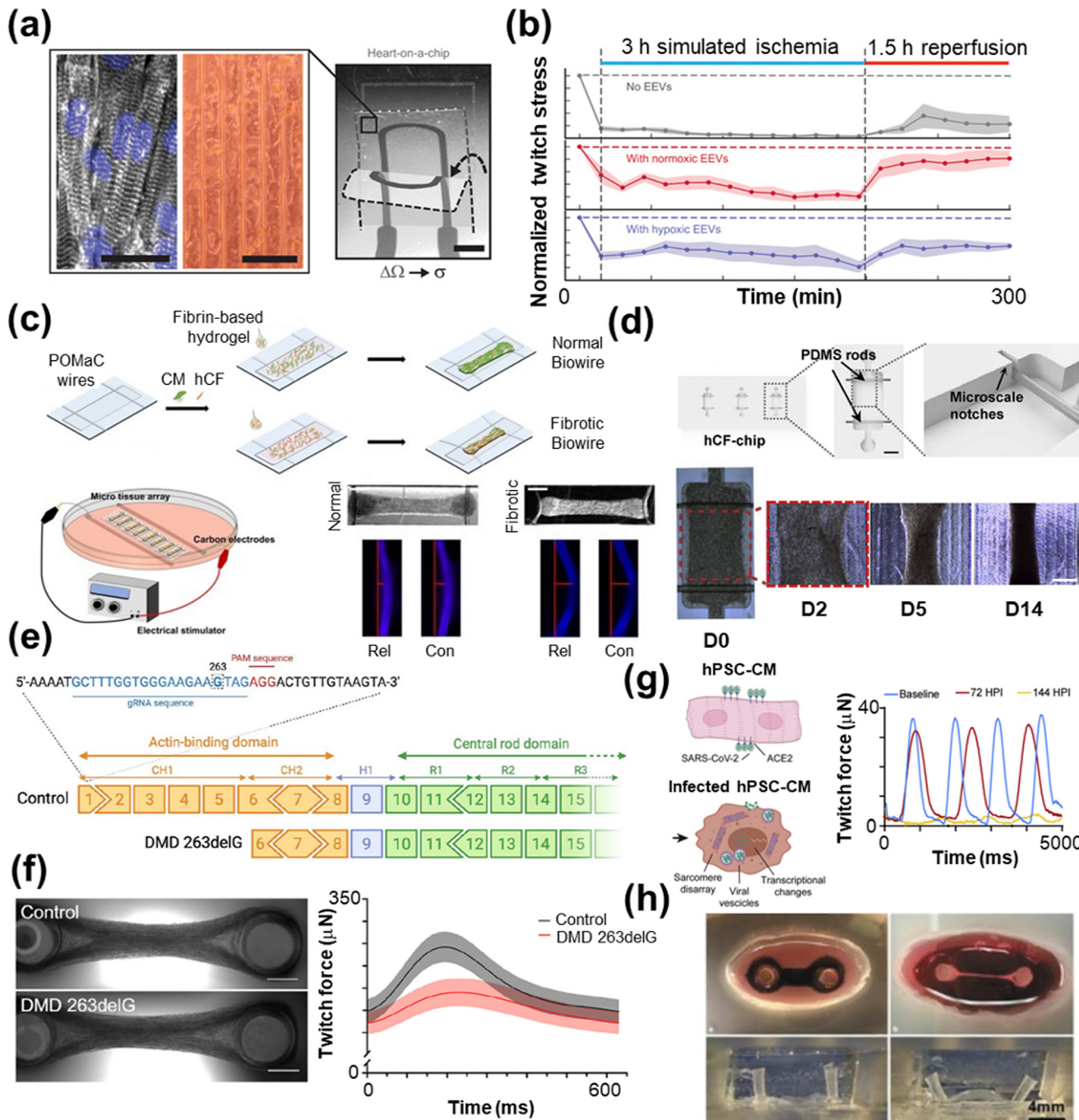


Fig. 4 Disease modeling with HoC systems. a) A MTF with an integrated strain sensor (scale bar = 2 mm) and microgroove template for cardiac tissue alignment (scale bars = 100 μ m) to monitor contractile behavior during ischemia-reperfusion. b) The resulting twitch force measurements post-extracellular vesicle treatment (reproduced from ref. 264 with permission from the American Association for the Advancement of Science). c) A Biowire model of cardiac fibrosis comprised of poly(octamethylene maleate (anhydride) citrate) (POMaC) wires are carbon electrodes for electrical conditioning (scale bar = 500 μ m) (reproduced from ref. 285 with permission from the American Chemical Society). d) A human cardiac fibroblast (hCF) chip with PDMS rods (scale bar = 1 mm) to support the tissue over the course of 14 days (scale bar = 500 μ m) (reproduced from ref. 286 with permission from Elsevier). e) An example gene editing process using the CRISPR/Cas9 technique, which is particularly useful for modeling genetic diseases. f) A two-pillar assay of control and tissue with Duchenne muscular dystrophy (DMD), which shows lower twitch force compared with control tissue (scale bar = 1 mm) (reprinted from ref. 288 under Creative Commons NonCommercial License 4.0 (CC BY-NC)). g) hPSC-CMs infected with SARS-CoV-2 show lower twitch force compared to the control 72 and 144 hours past infection (HPI) (reproduced from ref. 289 under Creative Commons License CC BY 4.0). h) Optical images of a biomimetic cardiac tissue model (BCTM) before loading cells (top left) and after establishing the engineered heart tissue (EHT) (top right). The system can mimic the end of systole (bottom left) and diastole (bottom right), as shown without EHT present (reproduced from ref. 290 with permission from Karver Publishers).

CMs displaying a protein profile that more closely resembled uninjured CMs. A key finding here is that the cardio-protective effects of EVs require their uptake by CMs, indicating an intracellular mechanism (Fig. 4a and b). Ellis *et al.* followed up this work with a new HoC system that places hiPSC-CMs and hiPSC-ECs in co-culture to better mimic the vasculature of the human heart. By analyzing three characteristic EV markers (CD9, CD63, and TSG101) at different points of the IR episode, the authors found that specific EV subpopulations are released at various stages of IR. Additionally, their system integrated on-board biosensors and EV lysis chamber based on standing acoustic waves (SAW), which enabled the analysis of the EV internal contents, particularly miR-1, miR-208b, and miR-499. Comparison of miRNA levels between the HoC system and clinical samples showed very good agreement at both ischemic and reperfusion stages, indicating the *in vitro* system could replicate some *in vivo* behavior.

As demonstrated by these reports, incorporating multiple cell types is crucial for recapitulating the native human physiology. Crucially, intercellular communication *via* EV release and other paracrine signaling processes has garnered tremendous interest for both improving the fundamental understanding of IRI and MI, as well as a potential target for early diagnosis and therapeutic intervention.^{265,278}

4.1.3 Cardiac fibrosis and remodeling. Closely related to MI, cardiac fibrosis describes the accumulation of ECM proteins within the cardiac interstitium, which results in increased stiffening of the heart muscle and reduced contractile function.^{279,280} Cardiac fibrosis is a reparative process associated with or a result of IRI, aging, diabetes, genetic cardiomyopathies, or metabolic syndrome. Fibrotic remodeling is often caused by a substantial loss of cardiomyocytes, for example during IRI. Because the heart has limited regenerative ability, the damaged tissue is replaced with collagen-rich scar tissue by myofibroblasts in an effort to maintain the structural integrity of the heart and prevent rupture. The alteration of the mechanical properties of the myocardium can lead to poor contractile force, poor electrical connection between myofibroblasts and CMs, arrhythmia, and reduced ejection fraction, which, as a result, can further lead to cardiac hypertrophy and accelerated decline in heart function.

Ugolini *et al.* developed a HoC system to study mechanical stimulation on CFs *in vitro*.¹⁴² Mechanical strain did have an effect on CF morphology, including cell area and preferential alignment of the cellular main axis orthogonal to the strain direction. An increase in CF proliferation was also reported for cells subject to 2% and 8% mechanical strain. Occhetta *et al.* leveraged a HoC platform with mechanical actuation to study the formation of cardiac scar tissue *in vitro*.²⁸¹ Using cardiac fibroblasts suspended within a fibrin hydrogel, the authors examined the effects of mechanical stimulation and exposure to the pro-fibrotic growth factor TGFβ-1 on scar tissue evolution. Hallmarks of cardiac wound-healing, include fibroblast proliferation, fibroblast to myofibroblast transition, matrix deposition, and tissue stiffening were seen.

TGFβ-1 was found to have a profound impact on fibroblast proliferation while mechanical stimulation (both with and without TGFβ-1) led to more pronounced transition from fibroblast to myofibroblast phenotype, which was also seen by Kong *et al.*²⁸² Interestingly, the increased level of myofibroblasts as a result of mechanical stimulation also coincided with a more homogeneous and stiffer tissue, likely attributable to the central role myofibroblasts play in ECM generation and deposition.

Given that cardiac fibrosis is intrinsically a multi-cellular process between CMs and (myo)fibroblasts, examples of HoC platforms typically contain multiple cell types to recapitulate this behavior. Lee *et al.* developed an HoC platform with cardiac microtissues that combined hESC-CMs and mesenchymal stem cells (MSCs) as a source of fibroblasts.²⁸³ Subjecting the microtissue to TGFβ-1 for two weeks resulted in irregular contraction patterns and myofibroblast differentiation, a result also seen by Sadeghi *et al.*²⁸⁴ Moreover, a thicker collagen layer was observed despite an overall decrease in microtissue diameter caused by CM apoptosis. Wang *et al.* also reported increased collagen density, impaired contractile function, increased tissue stiffness and passive tension, reduced active force, and increased excitation threshold in a HoC system that incorporated electrical stimulation and parallel rods to monitor contraction mechanics (Fig. 4c).²⁸⁵ Mastikhina *et al.* observed similar behavior after exposing cardiac microtissue to TGFβ-1 for three weeks (Fig. 4d).²⁸⁶ By incorporating parallel PDMS rods for mechanical characterization within the culture chamber, the authors were better able to quantify the degradation of contractile function. For instance, after 21 days of TGFβ-1 exposure, contraction force was approximately 7 times lower compared to the control. The authors also treated the tissue with the anti-fibrotic drug pirfenidone and measured a reduction in tissue stiffness. However, pirfenidone was not able to fully reverse tissue fibrosis. In a follow up report using a similar system, Mourad *et al.* showed fibrotic tissue possessed a genetic profile indicative of senescence.²⁸⁷ Treatment with a combination of the anti-senescence drugs dasatinib and quercetin led to improvements in active force and a downregulation of many senescence markers, such as tumor-suppressors CDKN1A and CDKN2A and cytokines such as interleukin-17A (IL-17A), IL-1 α , and macrophage inflammatory protein 1 α/β . However, no difference in collagen density or cell proliferation was seen after drug treatment.

4.1.4 Genetic and inherited cardiomyopathies. Cardiomyopathies can manifest from a variety of genetic mutations which cause dysfunctional energy metabolism, structural irregularities, or poor ionic homeostasis. Recent advances in gene editing along with the ability to obtain cardiac cells from patient-derived iPSCs have enabled researchers to study genetic and inherited cardiomyopathies in human relevant models.²⁹¹ While *in vitro* models, HoCs, and engineered cardiac tissue cannot fully recapitulate all of the intricacies of the human heart, they have led to

improvements in disease modeling and evaluation of potential treatments. Furthermore, the rarity of many genetic disorders make it difficult, if not impossible, to establish a large enough cohort for a clinical trial. This challenge could be overcome through the use of stem cell-derived cardiac cells and new gene editing methods. In this section, we highlight and discuss examples of various inherited and genetic cardiomyopathies that have been modeled with HoC and engineered heart tissue platforms.

4.1.4.1 Barth syndrome. Barth syndrome (BTHS) is a rare, but under-diagnosed, multi-system disorder that occurs as a result of mutations or deletions of the tafazzin gene (*TAZ*), which is responsible for the acylation of cardiolipin, a major phospholipid of the inner mitochondrial membrane.^{292–296} BTHS can cause a variety of cardiomyopathies, including prolonged QTc interval, hypertrophy, and arrhythmia, among others. Wang *et al.* developed a HoC model of mitochondrial cardiomyopathy using CMs derived from hiPSCs collected from two patients with BTHS.²⁹⁷ BTHS-hiPSC-CMs demonstrated impaired sarcomere assembly and lower twitch and peak systolic stress when grown on a thin film cantilever assay, both of which contributed to increased production of ROS. The authors also measured increased basal oxygen consumption rates in diseased tissue. Contractile function of the BTHS-hiPSC-CMs tissue could be restored, however, through transfection with *TAZ*-modified RNA or treatment with linoleic acid, a precursor of mature cardiolipin,²⁹⁸ thereby demonstrating the potential of *in vitro* HoC systems for combined disease modeling and therapeutic evaluation.

4.1.4.2 Hypertrophic cardiomyopathy (HCM). Hypertrophic cardiomyopathy is a common (≈ 1 in 500)²⁹⁹ genetic heart disease that can lead to arrhythmic sudden death, heart failure, and atrial fibrillation.³⁰⁰ The disease is often, but not exclusively, characterized by a remodeling of the left ventricle, particularly changes in wall thickness.^{301,302} A majority (70%) of the genetic mutations responsible for HCM occur in two sarcomeric genes, *MYH7* and myosin-binding protein C (*MYBPC3*).³⁰⁰ To better understand the mechanisms that link these genetic modifications with particular HCM phenotypes, Cohn *et al.* leveraged the CRISPR/Cas9 gene editing technique to introduce mutations in the *MYH7* and *MYBPC3* genes. Using a two-pillar assay, they found the modified cardiac tissue exhibited altered Ca^{2+} transients, prolonged relaxation times, and increased twitch force and resting tension.³⁰³ The observed hypercontractile behavior was reversible to some degree through treatment with verapamil, a Ca^{2+} channel blocker, or blebbistatin, a direct myosin inhibitor, although only blebbistatin lowered both twitch force and resting tension. In the same year, Zhao *et al.* built upon their previously reported “Biowire” platform³⁰⁴ to study phenotypic differences between hypertensive patients with and without left ventricular hypertrophy (LVH).³⁰⁵ The Biowire II platform enabled electrical conditioning and culturing of hiPSC-CMs obtained from patients for up to 8 months. Engineered tissue from patients affected with hypertension and LVH consistently

showed an upregulation of genes associated with cardiac hypertrophy and heart failure, as well as an inability to generate a contractile active force after 8 months. In another report, Cashman *et al.* sought to better understand the role of the *BRAF* gene in patients with cardio-facio-cutaneous syndrome (CFCS) that show evidence of HCM.³⁰⁶ Using patient-derived hiPSC-CMs and cardiac stromal cells cultured on PDMS pillars, the authors reported a larger tissue cross-section, accelerated twitch dynamics, and higher levels of the hypertrophic marker atrial natriuretic peptide (ANP) in *BRAF*-mutant tissue, which is in agreement with other reports of hypertrophic tissue.^{307,308} The E99K mutation in the *ACTC1* gene (encodes α -cardiac actin) was studied using patient-derived hiPSC-CMs from a patient displaying HCM and left ventricular non-compaction (LVNC) along with the patient's two sons (one non-carrier and one carrier with a normal ECG).³⁰⁹ Interestingly, while the hiPSC-CMs derived from the father showed numerous pathogenic phenotypes, most were not observed in hiPSC-CMs derived from the two sons, even those with the E99K mutation. The common pathogenic phenotype was arrhythmia as determined with a two-pillar assay. The distinct differences between diseased tissues highlights the complex interplay between a particular mutation, background genetics, and age-dependence.

4.1.4.3 Duchenne muscular dystrophy (DMD). Duchenne muscular dystrophy is an X-chromosome linked, muscle-wasting disorder that is prevalent in less than 100 in 1 000 000 males and less than 1 in 1 000 000 females.^{310–312} Mutations to the *DMD* gene alter the encoding of muscular dystrophin, making the muscles more susceptible to damage and subsequent loss of functionality. As a result, cardiomyopathy can arise and eventually lead to heart failure.³¹⁰ Bremner *et al.* leveraged CRISPR/Cas9 to introduce a mutation in the *DMD* gene (DMD 263delG) of hiPSCs derived from a healthy patient to delineate genotype-phenotype correlations (Fig. 4e and f).²⁸⁸ After differentiation to hiPSC-CMs, a two pillar HoC platform was used to study the contractile behavior of the engineered microtissue. The modified tissue showed a dysregulation of biological processes related to cardiac muscle development, contraction, membrane potential regulation, Ca^{2+} handling, and ECM organization. Measurements of contractile function confirmed lower twitch force, beating irregularities, and slower kinetics in modified tissue, while fluorescence imaging revealed elevated cytosolic Ca^{2+} levels and prolonged Ca^{2+} transients. Macadangdang *et al.* also saw similar behavior that was more pronounced in hiPSC-CMs subjected to a combinatorial maturation (ComboMat) procedure compared to immature hiPSC-CMs, emphasizing the importance of cardiac maturation in *in vitro* HoC systems.³¹³

4.1.4.4 Dilated cardiomyopathy (DCM). Dilated cardiomyopathy is one of the most common cardiomyopathies in the world.³¹⁴ This disorder is typically characterized by the dilation of the left or both ventricle(s) and systolic dysfunction that cannot be attributed to pressure-volume overload or coronary artery disease.^{314,315} Alterations of genes that encode

cytoskeletal and sarcomeric proteins are a common cause of inherited DCM.³¹⁴ Examples of causative genes are *TTN* (encodes titin), *DMD* (encodes dystrophin), *LMNA* (encodes lamin A/C), *MYH7* (encodes β -myosin heavy chain), *RBM20* (encodes RNA binding motif protein 20), and *TNNI1* (encodes cardiac troponin 1), among others, especially those implicated in HCM.^{177,314-316} The effect of a missense mutation S635A in the *RBM20* gene was examined by Streckfuss-Bomeke *et al.* using DCM patient-derived hiPSC-CMs.³¹⁷ Although no differences in cell surface area were seen, the modified tissue showed a markedly irregular distribution of the sarcomeric protein α -actinin along with dysfunctional Ca^{2+} cycling. As a result, the modified tissue demonstrated lower contractile force and decreased passive stress compared to control samples. The *RBM20* mutation also led to preferential retention of the larger and more elastic *TTN* protein isoform N2BA as opposed to smaller and stiffer N2B isoform, which is predominant in a healthy heart³¹⁸ while higher N2BA content has been reported in failing hearts.^{319,320}

Mutations of the *TNNT2* gene also caused sarcomeric irregularities and contractile dysfunction.³²¹ Dai *et al.* investigated a particular mutation, R173W, and confirmed poor sarcomeric organization and contractile dysfunction using a two-pillar assay.³²² The mutated troponin T (TnT) showed reduced binding with tropomyosin (Tm), which could be the cause for poor sarcomeric organization. Activation of AMP-activated protein kinase (AMPK), which interacts with myosin heavy chain 7, using A-769662 led to improved contractile function and sarcomeric alignment, suggesting a potential therapeutic pathway for improving DCM recovery.

In addition to sarcomeric and cytoskeletal genes, Wauchop *et al.* recently reported on the impact of mutations to the sodium voltage-gated channel alpha subunit 5 (*SCN5A*) gene, which is responsible for encoding the cardiac sodium channel (Nav1.5).³²³ Patient-derived hiPSC-CMs with the R222Q missense mutation to *SCN5A* were first seeded as isolated single cells or cardiac sheets, although both failed to display the electrophysiological characteristics associated with the R222Q mutation. This is most likely due to tissue immaturity as the R222Q mutation is only expressed in adult isoforms of *SCN5A*.³²⁴ To promote hiPSC-CM maturation, the tissue was seeded in the previously reported Biowire platform²⁸⁵ for long-term culture. Expression of the adult variant of *SCN5A* was $\approx 90\%$ in mature tissue, and as a result, the disease phenotype (*i.e.* contractile dysfunction, dilation, sarcomeric irregularity) was seen, which again highlights the importance of tissue maturity in *in vitro* disease modeling.

4.1.4.5 Pompe disease. Pompe disease is a severe autosomal recessive disorder that typically, although not exclusively, presents itself during infancy and is caused by the loss of acid α -glucosidase (GAA).³²⁵ GAA is the enzyme responsible for converting glycogen into free glucose within lysosomes. Accumulation of glycogen can result in muscle weakness, respiratory issues, and cardiac hypertrophy. To study the effect of Pompe disease on cardiac function, Raval *et al.* used two patient-derived hiPSC-CM cell lines, each possessing different

mutations in the *GAA* gene (del ex18/del ex18: deletion of exon 18 and 1441delT/2237G>A: deletion of a T nucleotide at *GAA* cDNA position 1441 and G to A transition at *GAA* cDNA position 2237).³²⁶ Accumulation of glycogen within the lysosome was confirmed by electron microscopy. Contractile function was evaluated using a two-pillar assay, and the del ex18/del ex18 hiPSC-CMs demonstrated accelerated kinetics and reduced paced peak contractile force. Nonetheless, contractile function was largely similar between diseased and control tissues.

4.1.4.6 Friedreich's ataxia. Friedreich's ataxia is an autosomal recessive disease that is caused by a mutation in the *FXN* gene, which suppresses the expression of frataxin, a small mitochondrial protein.³²⁷ Cardiac dysfunction is the most attributed cause of death in patients with Friedreich's ataxia.³²⁸ Using two patient-derived hiPSC-CM lines and human fibroblasts, Wong *et al.* utilized the two-pillar PDMS chip to study contractile function in engineered cardiac tissues.³²⁹ Expression of *FXN* was confirmed to be about 70% lower at the transcriptional level and 40% to 60% lower at the protein level in diseased tissue compared to the control. Force generation was less than half in diseased tissue and contraction kinetic were significantly slower in diseased tissue compared to control samples. Contractile dysfunction was largely reversed, however, after transduction with lentiviral-FXN.

4.1.4.7 Hypoplastic right heart syndrome (HRHS). Hypoplastic right heart syndrome is characterized by an underdeveloped right ventricle and can be accompanied by a defect in the atrial septum.³³⁰ Using a two-pillar PDMS chip, Lam *et al.* seeded patient-derived hiPSC-CMs with human fibroblasts to study contractile function.³³¹ The tissue was subjected to 1 Hz, 1.5 Hz, and 2 Hz electrical pacing and showed significantly reduced contractile force compared to the control samples. The contraction kinetics also remained constant in diseased tissue over the course of 2 weeks, whereas control samples showed a gradual acceleration as the cells matured. Furthermore, Ca^{2+} dynamics and electrophysiology were comparable between normal and diseased tissue, suggesting the calcium handling remained intact. Diseased tissue was found to downregulate gene transcripts involved in cardiac maturation and development (natriuretic peptide B (*NPPB*), four and a half LIM domains 2 (*FHL2*), PDZ and LIM domain 3 (*PDLIM3*)) and cardiac contraction (*MYH7*, *MYL2*, *etc.*), suggesting abnormal and immature cardiac development.

4.1.4.8 Catecholaminergic polymorphic ventricular tachycardia (CPVT). Catecholaminergic polymorphic ventricular tachycardia is an exercise- or emotional-induced polymorphic ventricular tachycardia that occurs in the absence of structural abnormalities.^{332,333} Many incidences of CPVT are caused by mutations to the *RyR2* or *CASQ2* genes, which encode proteins that are responsible for Ca^{2+} handling and transport. Using engineered heart tissue on a thin film cantilever array and Ca^{2+} imaging, Park *et al.* demonstrated hallmarks of CPVT with patient-derived iPSC-CMs, including re-entrant arrhythmia after electrical pacing and catecholamine (isoproterenol)

administration.³³⁴ Furthermore, the authors were able to show the phosphorylation of ryanodine receptor type 2 (RYR2) at the serine-2814 position by Ca^{2+} /calmodulin-dependent protein kinase II (CaMKII) is needed to realize the arrhythmic potential in CPVT patient-derived cardiac tissue.

4.1.5 Severe acute respiratory coronavirus-2 (SARS-CoV-2) and cardiac dysfunction. Severe acute respiratory coronavirus-2 is the virus responsible for causing coronavirus disease 2019 (COVID-19), which was first reported in Wuhan, China in late 2019.¹⁸¹ Since then, the World Health Organization has reported almost 770 million confirmed cases and nearly 7 million deaths worldwide as of August 2023, although the actual count could be much higher.³³⁵ Although COVID-19 is predominantly a respiratory disease, it has also been implicated in various cardiovascular complications, such as arrhythmia, myocardial infarction, and myocarditis.^{16,336,337} SARS-CoV-2 viral particles were even found in cardiac tissue obtained from a child who died following COVID-19 early on in the pandemic.³³⁸ Nonetheless, the mechanism(s) by which COVID-19 causes cardiovascular dysfunction, be it directly through myocardial infection or indirectly through virus-associated systemic inflammation, is not entirely understood. To improve upon this, Bailey *et al.* developed an engineered heart tissue model with hiPSC-derived cardiomyocytes, fibroblasts, and/or macrophages to study COVID-19 myocarditis.³³⁹ The two-pillar PDMS system showed infected cardiomyocytes primarily at the periphery of the tissue along with macrophage accumulation in the infected regions. Using a two-pillar assay, the infected tissue also demonstrated reduced contractile function and slower kinetics, which resulted from sarcomere loss and reduced troponin T expression. The authors tested two therapeutic agents, a TANK-binding kinase 1 (TBK1) inhibitor, which blocks viral nucleic acid sensing pathways, and remdesivir, which inhibits viral replication. While both remdesivir and TBK1 inhibition prevented reductions in *TNNT2* and *MYH7* mRNA expression, only remdesivir could prevent sarcomere loss and cell death suggesting cardiomyocyte infection as a root cause of disease rather than inflammation. Marchiano *et al.* also generated engineered cardiac tissue from hPSCs to study SARS-CoV-2 infection (Fig. 4g).²⁸⁹ CMs were found to express ACE2, making them susceptible to viral entry and infection, whereas smooth muscle cells did not. Two multiplicities of infection (MOI) were studied, 0.1 and 5. Both MOIs led to viral infection and replication within the CMs, even causing cessation of beating and cell death after 48 h at MOI = 5. Using a multi-electrode array (MEA), the authors noted a decline in the electrophysiological properties of the infected cells, including a reduced beating rate, lower depolarization spike amplitude, and decreased electrical conduction velocity. Furthermore, using a two-pillar PDMS chip with an embedded magnetic sensor, a degradation in the contractile properties could be seen beginning at 48 h post-infection with the twitch force falling to near 0 after 6 days, which could be attributed to decreased CM density and a loss of sarcomeric organization. These results again suggest that SARS-CoV-2 can directly cause heart damage independent of inflammation. A similar decline in contractile

function was reported in a pre-print paper from Xing *et al.* who developed a HoC platform that mimicked the myocardium and vasculature based on their previously reported InVADE platform.^{340,341} However, effects to contractile function could only be seen after incorporating peripheral blood mononuclear cells (PBMC) into the chip, which resulted in increased production of cytokines and chemokines and higher levels of circulating cell-free mitochondrial DNA. Interestingly, these effects were largely reversible upon treatment with EEVs, indicating a potential therapeutic pathway to minimize inflammation in patients with SARS-CoV-2 induced myocarditis.

4.1.6 Mechanical stress. A distinct advantage of *in vitro* HoC systems over conventional 2D static cell cultures is the ability to incorporate mechanical stimulation within the cellular microenvironment. In the case of stem cell-derived CMs, mechanical stimulation has been demonstrated as a means to advance maturation,^{163,342} which has remained a fundamental challenge in their usage for disease modeling and drug testing. On-chip conditioning of CMs with mechanical stimulation aims to mimic the various mechanical forces experienced *in vivo*, including shear stress from blood flow, cyclic strain during beating, stretching due to blood pressure changes, and forces due to changes in elastic modulus over time.^{124,141,204} While mechanical stimulation promotes cardiac maturation, it has also been implicated in various cardiopathologies.³⁴³ For instance, Rogers *et al.* utilized a two pillar design capable of mimicking the rhythmic pressure and volume changes of the heart to study the effects of mechanical stimuli (Fig. 4h).^{178,344} Both volume and pressure overload resulted in structural changes to the tissue, altered gene expression, and oxidative stress. Pressure overload conditions in particular were associated with concentric hypertrophy and increased fibrotic signatures, while tissue subjected to volume overload showed myocyte thinning and matrix degradation. Leonard *et al.* also measured changes to gene expression in hiPSC-derived cardiac microtissues subject to varying degrees of afterload cultured in an adjustable two post system.¹⁶³ While low levels of afterload (*i.e.*, $\approx 0.1 \mu\text{N} \mu\text{m}^{-1}$) improved maturation, higher levels of afterload (*i.e.*, $\approx 10 \mu\text{N} \mu\text{m}^{-1}$) resulted in upregulation of hypertrophic markers, including ANP, brain natriuretic peptide (BNP), and α -skeletal actin (ACTA1), metabolic changes from fatty acid utilization (mature-like) to glycolysis (fetal-like), and increased expression of fibrosis-associated growth factors, similar to other reports.^{176,345} Despite changes in gene expression, the authors noted the tissue stiffness remained unchanged despite the increase in fibrosis markers.

4.2 Heart-on-a-chip for drug screening and cardiotoxicity assessment

HoC systems have tremendous potential in drug development and cardiotoxicity applications, which are currently dominated by animal studies. While animal models have been instrumental for evaluating therapeutics, there remains

significant disagreement between results of animal studies and clinical trials^{346–349} prompting some to call into question their use.³⁵⁰ Moreover, animal models contribute millions of US dollars and years of time to the already significant cost and time associated with drug development,^{10,351} which takes around 1 billion USD and 12 years per drug.^{352,353} Nonetheless, animal studies oftentimes fail to adequately capture human (patho)physiology and pharmacology, leading to poor translation to clinical trials³⁴⁹ or removal from the market due to safety concerns.³⁵⁴ Because of this poor translation, policies surrounding animal testing are beginning to change, as evidenced, for example, by the US Food and Drug Administration (FDA) recently lifting the

requirement of animal testing for new drugs.³⁵⁵ Traditional 2D cell culture, while inexpensive and well-established, is also largely limited in its capacity to recreate the disease phenotypes seen *in vivo*, likely due to the mechanical properties of culture plates and altered pharmacokinetics and pharmacodynamics. The use of *in vitro* human microphysiological models is a viable alternative to conventional 2D cell culture due to their potential for automation, human relevance, and improved control over experimental conditions. Furthermore, the ability to obtain patient-derived PSCs has opened the door to increasingly personalized disease treatment, while advances in microfabrication and automation have improved throughput

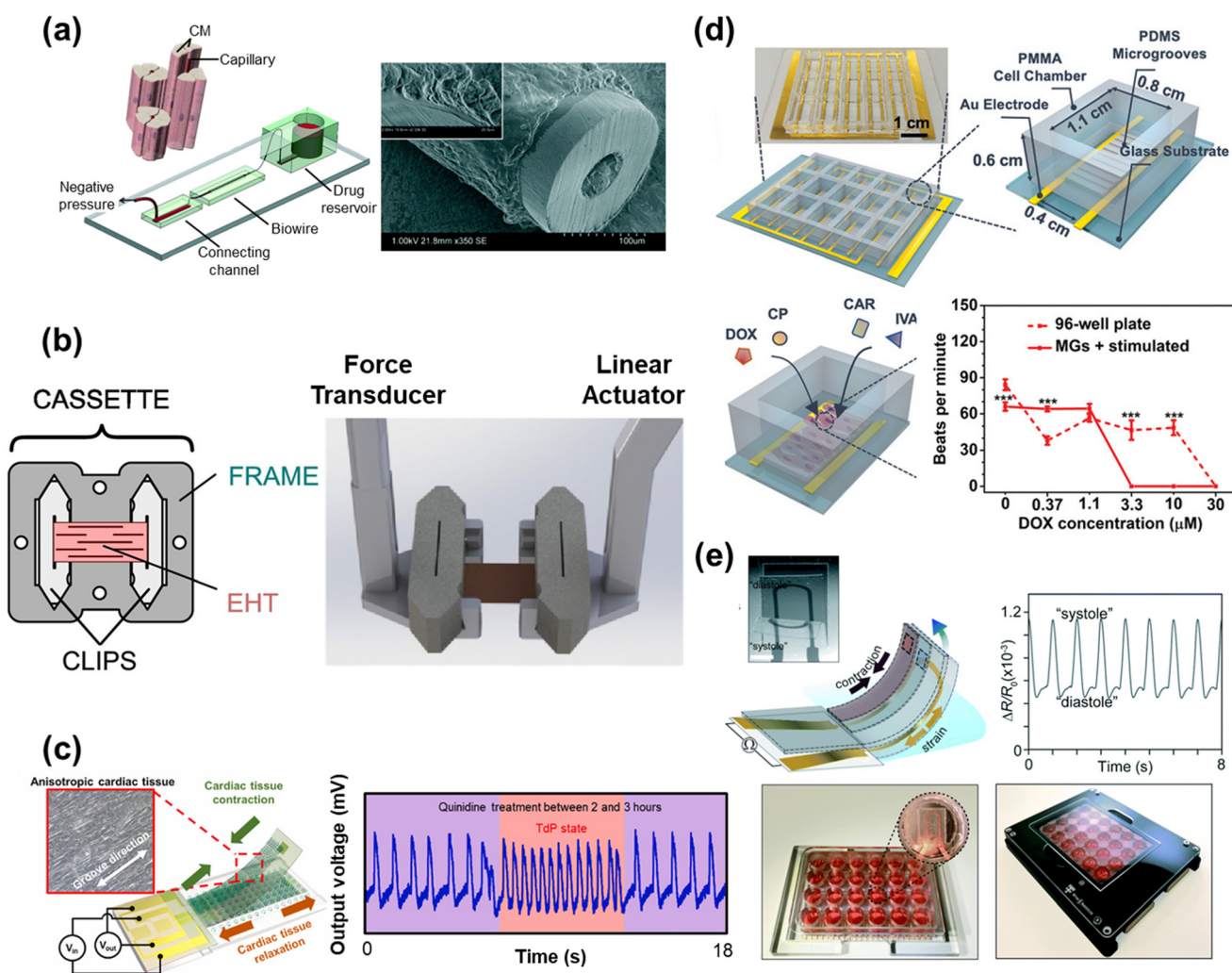


Fig. 5 Drug efficacy and toxicity assessment with HoC systems. a) A perfusable Biowire platform to mimic drug diffusion through the capillaries and muscle tissue, used here for studying the effect of nitric oxide on muscle tissue (reproduced from ref. 304 with permission from the Royal Society of Chemistry). b) A custom cardiac tissue holder capable of mechanical characterization and adjustable loading (reproduced from ref. 344 with permission from Elsevier). c) A flexible cantilever with an integrated strain sensor and patterned microgrooves to promote tissue alignment. The resulting signal output after treatment with quinidine shows a torsades de pointes (TdP) state (reproduced from ref. 182 with permission from Elsevier). d) A HoC platform with patterned microgrooves (MGs) and electrical stimulation capabilities for evaluating the cardiotoxicity of antineoplastics (doxorubicin (DOX) and cyclophosphamide (CP)) and potential drugs to ameliorate cardiotoxicity (ivabradine (IVA) and carbachol (CAR)) (reproduced from ref. 378 with permission from John Wiley and Sons). e) A flexible cantilever with an integrated strain sensor for monitoring contractile behavior as part of a larger array for high-throughput drug toxicity screening (reproduced from ref. 379 with permission from the Royal Society of Chemistry).

tremendously, potentially leading to rapid screening and characterization of drug–drug interactions. In the following section, we discuss in more detail reports of HoC systems for drug/therapeutic screening and cardiotoxicity assessment.

4.2.1 Evaluation of drug and therapeutic efficacy. The high costs and long timeline (10+ years)³⁵⁶ currently required to bring a drug to market has prompted the search for alternatives to animal testing. Furthermore, many drugs fail to make it through clinical trials despite the large resource requirements, with only a $\approx 10\%$ success rate. To reduce time-to-market and improve success rate, HoC systems could provide a valuable link between pre-clinical animal studies and clinical trials or even serve as a standalone replacement to animal studies all together. One of the most popular drugs tested with HoC systems is the non-selective β -adrenergic receptor agonist isoproterenol,^{143–145,155,160,168,172,181,224,225,357–368} which can help treat bradycardia. Isoproterenol is similar in structure and function to epinephrine, which has also been evaluated using HoC systems.^{369,370} The response to either drug *in vitro* has been similar to clinical data, which show responses in the range of 5 nM to 800 nM.^{371–375}

Using the Biowire platform, Xiao *et al.* studied the effect of sodium nitroprusside (SNP) perfusion on cardiac tissue (Fig. 5a).³⁰⁴ The system was designed such that the cardiac tissue grows circumferentially around a suspended lumen, which mimics the capillaries found in the native myocardium and helped to align the CMs. SNP, a nitric oxide (NO) donor, was perfused through the lumen and NO levels in the surrounding medium reached 100 μM , which is higher than physiological levels *in vivo*. NO is a vasodilator released by endothelial cells in the native myocardium and plays a crucial role cardiovascular disease progression and prevention.³⁷⁶ In the Biowire, NO was found to decrease beating frequency of the cardiac tissue due to a degradation of the myofibrillar cytoskeleton as determined by α -actinin fluorescent imaging. In another iteration of the Biowire platform (Biowire II), this time with the tissue grown perpendicular to and anchored to two suspended wires, Wang *et al.* evaluated the efficacy of three drugs used to treat angiotensin II (Ang II)-induced cardiomyopathy, losartan, relaxin, and saracatinib.³⁷⁷ Ang II treatment of the microtissue showed various hallmarks of disease, including increased passive tension, an acute positive inotropic response followed by a chronic negative inotropic response, and an increase of the excitation threshold. After 2 weeks of treatment, all 3 of the drugs reduced passive tension, although saracatinib also reduced the active force, indicating tissue degradation. Relaxin appeared to be the most effective at reducing passive tension, increasing active force, and rescuing the electrophysiological properties of the tissue, all at a dose determined to be safe from toxicity tests conducted on the same system.

For the treatment of HCM, Sewanan *et al.* and Prondzynski *et al.* examined two potential therapeutic candidates, mavacamten (a β -blocker) and diltiazem (a Ca^{2+} -channel blocker), respectively.^{380,381} The Campbell group

developed a custom tissue holder that was capable of on-board mechanical characterization and adjustable loading (Fig. 5b).^{376,344} Using this platform, mavacamten treatment was shown to improve diastolic tissue stiffness and relaxation time, as well as reduce the myocardial workload and power output, which would help lower the energetic cost of contraction caused by HCM.^{382,383} Prondzynski *et al.* also observed a reduction in absolute force, relaxation time, and APD after treatment with 3 μM diltiazem using a two-pillar HoC system. Here, iPSC-CMs were derived from a patient with HCM that exhibited a rare *ACTN2* mutation. Treatment with diltiazem resulted in more drastic changes in the HCM patient-derived iPSC-CMs compared to the isogenic control prepared using CRISPR/Cas9. Motivated by these improvements, diltiazem was prescribed to the son and sister of the original patient, both of whom carried the same mutation. Diltiazem was shown to reduce the QT interval from 460 ms to 387 ms in the son and from 477 ms to 439 ms in the sister, implicating diltiazem as a candidate for future clinical treatment of HCM.

4.2.2 Cardiotoxicity assessment. The most well-known families of drugs to induce cardiotoxicity are antineoplastics (e.g., anthracyclines, VEGF- and kinase inhibitors, taxanes),³⁸⁴ antibiotics (e.g., chloroquine), antiarrhythmics (e.g., quinidine), opiates (e.g., methadone), Ca^{2+} channel blockers (e.g., verapamil), and antihistamines (e.g., astemizole).³⁵⁶ Even after a drug makes it to market, recalls are not uncommon. Adverse cardiovascular effects are the primary cause for nearly 20% of the drugs that are recalled from the market in the US.³⁸⁵ Some of the most common cardiovascular complications include arrhythmia, myocyte toxicity, myocardial infarction, thrombosis, low blood pressure, and coronary artery disorders.^{356,384–387} Oftentimes, these detrimental effects are not seen in preclinical animal testing.¹⁰ Perhaps one of the most high-profile recent examples of this is the 2004 recall of rofecoxib, which is a non-steroidal anti-inflammatory drug (NSAID).^{10,388,389} A study by Graham *et al.* estimated that rofecoxib caused between 88 000 and 140 000 cases of serious coronary heart disease, including acute myocardial infarction requiring admission and sudden cardiac death, before being voluntarily pulled from the market in late 2004.³⁹⁰

4.2.2.1 Ion-channel blockers. Numerous HoC platforms with different analytical capabilities have been developed to study drug-induced cardiotoxic side effects.¹²⁶ One of the most commonly observed side effects is arrhythmia, including bradycardia, atrial fibrillation, long QT syndrome, and torsades de pointes (TdP). Many of the proarrhythmic drugs block one or more of the ion channels found in the heart (K^+ , Na^+ , Ca^{2+}), making HoC systems well-suited to study their effects. Examples include verapamil (a Ca^{2+} channel blocker),^{168,180,182,183,360,365,366,369,391–394} quinidine (Na^+ -channel blocker),^{182,359,392,395} E-4031 (hERG K^+ -channel blocker),^{180,359,360,363,396–398} sotalol (K^+ -channel blocker),^{305,361,396,397} nifedipine (a Ca^{2+} -channel blocker),³⁹⁶ flecainide (Na^+ -channel blocker),^{362,397,398} tetrodotoxin (Na^+ -channel blocker),³⁹⁸ ouabain (Na^+/K^+ -ATPase

blocker),³⁵⁹ ATX-II (Na^+ -channel blocker),³⁵⁹ and dofetilide (K^+ -channel blocker).^{359,395} The Lee group has developed various iterations of cantilever-based sensors to monitor real-time changes in contractile function after exposing cardiac tissue to different concentrations of drugs known to be cardiotoxic, including quinidine, E-4031, and verapamil.^{180,182,183,391,392} The cell culture surface is patterned with a series of microgrooves to promote tissue anisotropy and maturation (as discussed in section 4.1.1) and contractile function is characterized by measuring cantilever displacement (Fig. 5c). Using this technology, the group measured various forms of arrhythmia, including prolonged beating duration caused by E-4031,^{180,359,360,363,396–398} decreased contraction force and bradycardia after verapamil and lidocaine exposure,^{180,182,183,391,392} early after depolarization (EAD) and TdP at high quinidine levels.^{182,392} Given that arrhythmia is typically detected using electrocardiogram (ECG) and arises due to a dysfunctional conduction system, electrical measurements are well-suited to record drug-induced changes in field potential duration (FPD), which is analogous to the QT interval used in the clinic.^{399,400} Visone *et al.* developed a microelectrode chip that was benchmarked with three drugs, aspirin, sotalol, and verapamil.³⁹³ While aspirin showed no significant changes in FPD, both sotalol (a K^+ channel blocker) and verapamil showed substantial changes, in agreement with other reports.³⁹⁴ Interestingly, after verapamil treatment, the human tissue showed a dose-dependent decrease in FPD while the rat model showed a dose-dependent increase, reiterating the importance of using human-relevant *in vitro* models for assessing drug toxicity.

4.2.2.2 Antineoplastics. Certain antineoplastic therapies, while effective for cancer treatment, have also been implicated as cardiotoxic.³⁸⁴ Various chemotherapeutics have been studied using HoC systems, including linsitinib,³⁵⁸ oxaliplatin,⁴⁰¹ and cyclophosphamide (CP).⁴⁰² One of the most widely used, clinically-approved, anti-cancer drugs is the anthracycline, doxorubicin (DOX).³⁸⁴ Anthracycline-induced cardiotoxicity is likely caused by an increase in the generation of reactive oxygen and nitrogen species (ROS/RNS).⁴⁰³ Given its well-established cardiotoxicity, many HoC reports demonstrate doxorubicin as a proof-of-concept.^{145,162,166,236,378,394,401,404} An example from Ren *et al.* reported the design of a HoC platform that combined electrical stimulation with patterned PDMS microgrooves to promote tissue maturation (Fig. 5d).³⁷⁸ The device was used to study the effects of DOX and CP, in addition to two potentially cardioprotective supplements, ivabradine (IVA) and carbachol (CAR). Both DOX and CP appeared to be chronotropic and cytotoxic, although only DOX caused a dose-dependent decrease in cell viability. When tested with DOX, both CAR and IVA fully restored cell viability and appeared to reduce the chronotropic effects of DOX. HoC systems have also been used to evaluate the cardiotoxicity of phytochemicals with potential anti-cancer properties, such as sulforaphane^{405,406} and thapsigargin,^{306,407,408} as well as kinase inhibitors,⁴⁰⁸ which have been widely explored as chemotherapeutics.⁴⁰⁹ The cardiotoxic effects of a range of kinase-inhibitors, specifically

microtubule affinity regulating kinase (MARK) and checkpoint kinase (Chk), were characterized using an impedimetric assay.⁴¹⁰ Two MARK-inhibitors synthesized by the authors caused dose-dependent decrease in beat amplitude also seen in tests with well-established microtubule targeting drugs, paclitaxel, vinblastine, nocodazole, and colchicine. The decrease in beat amplitude was determined not to be a result of cytotoxicity. Similar effects were also seen in all but two of the nine Chk-inhibitors tested with the impedimetric assay, although the half-maximal inhibitory concentration (IC_{50}) varied more than two orders of magnitude across samples.

4.2.2.3 Antimicrobials. Antimicrobial drugs, including antibiotics, antivirals, and antifungals, have saved countless lives since their introduction in the 20th century, perhaps even earlier.⁴¹¹ The cardiotoxicity of a selection of antimicrobial drugs, including ketoconazole,⁴¹² and azithromycin,⁴¹³ has been evaluated using HoC systems. Of particular relevance currently are the cardiotoxic effects of various treatments for COVID-19. Xu *et al.* investigated 4 repurposed antiviral drugs, namely remdesivir, apilimod, ritonavir, and lopinavir, using a human EHT model.⁴¹⁴ All 4 were found to affect Ca^{2+} transients and contractile function while causing broad transcriptional changes in the EHTs, although apilimod and remdesivir showed a far more altered transcriptome compared to ritonavir and lopinavir. Over 2000 potential protective drugs were also screened to reduce the remdesivir-induced cardiotoxicity. Astaxanthin was found to be an excellent protectant independent of its antioxidant activity, largely restoring cardiac function when used in tandem with remdesivir. The use of azithromycin (AZM), a macrolide antibiotic, and hydroxychloroquine (HCQ), an antimalarial drug, separately or combined, was also a widely used treatment for COVID-19 (ref. 415) despite a lack of evidence supporting any clinical benefit⁴¹⁶ and evidence suggesting they are cardiotoxic.^{417,418} Both Wong *et al.*⁴¹⁹ and Charrez *et al.*⁴¹³ validated the cardiotoxic properties of HCQ and AZM separately and jointly in human EHT systems. Each concluded HCQ and AZM have proarrhythmic effects and increase the action potential duration (APD), which is analogous to an increase in QT (long QT syndrome). In the context of a worldwide pandemic, like COVID-19, where time is crucial, HoC systems could be especially useful for rapid cardiotoxicity assessment of drugs under consideration for emergency use authorization.

4.2.2.4 Nanomaterials and other drugs. The cardiotoxicity of various other classes of drugs and nanomaterials has been studied using HoC systems, including antihistamines (e.g. terfenadine,^{172,378} astemizole,³⁹³ and fexofenadine¹⁷²), anesthetics^{361,371,393,397} steroids (e.g. dexamethasone¹⁴⁴ and hydrocortisone⁴¹²), anti-depressants⁴²⁰ pain relievers (e.g. acetaminophen^{166,412} and diclofenac⁴¹²), myosin modulators (e.g. blebbistatin^{363,364} and omecamtiv mecarbil³⁶⁴), anti-fibrotics²⁸⁶) beta blockers,^{181,362} prokinetics,³⁹⁸ phosphodiesterase inhibitors,⁴²¹ hormones,¹⁴⁴ and nanomaterials.^{379,422} Although this is a small sampling of the many therapeutics in pre-clinical studies, clinical trials, and on

the market, it highlights the broad range of drug types that could potentially be studied using HoC systems.

4.2.2.5 Medium- and high-throughput screening. Perhaps one of the most advantageous properties of *in vitro* HoC platforms is the potential for higher throughput, (semi-)automated, evaluation of large libraries of therapeutics. The result is highly robust datasets with redundant measurements taken in nearly identical environmental conditions. Pointon *et al.* developed an improved throughput workflow combining timelapses with automated image analysis to evaluate 29 inotropic and 13 non-inotropic compounds using 3D engineered cardiac microtissues.⁴²³ The system was capable of distinguishing between positive and negative inotropes and detecting contraction changes with 80% sensitivity and 91% specificity. Similar results were seen in a separate study from Ando *et al.*, where they evaluated a total of 60 drugs, 57 of which are known to be torsadogenic, from 4 distinct drug families using a microelectrode array (MEA) to measure changes in field potential duration.⁴²⁴ The developed assay was able to categorize drugs based on torsadogenic risk and when compared to the CredibleMeds database, yielded a sensitivity of 81%, specificity of 87%, and accuracy of 83%. A two-tiered assay was developed by Keung *et al.* to examine inotropic behavior in human ventricular-like cardiac tissue strips (hvCTS) and human ventricular-like organoid chambers (hvCOC).⁴²⁵ 25 drugs were screened, first with hvCTS to determine any negative inotropic response, then with hvCOC

to determine any positive inotropic response. The assay demonstrated accuracies of 100%, 86%, and 80% for negative, positive, and null inotropic responses, respectively. A 24-well plate-style HoC platform was developed by Lind *et al.*, which contained microgrooved PDMS cantilevers with integrated electronic strain gauges for on-board quantification of twitch stress and beat rate (Fig. 5e).⁴²⁶ The system was benchmarked by measuring the dose-dependent response to 12 distinct cardiac drugs. Furthermore, the open-well format allowed for the use of Transwell® inserts to model the pharmacological response to drug diffusing through an endothelial barrier, which was found to prolong the time to total cardiac failure compared to a direct exposure.

5 Heart-on-a-chip as part of multi-organ-on-a-chip platforms

Up to this point, we have discussed the heart and HoC systems as an isolated organ. However, in reality the human body exists as a complex interplay between its various organs. An advantage of organ-on-a-chip platforms is the ability to integrate two or more organs to study a particular organ-organ coupling or create a “body-on-a-chip”.⁴²⁷ This is especially relevant for drug and therapeutic testing, where a detailed understanding of the pharmacokinetics and pharmacodynamics could lead to significant improvements in drug development and clinical trials.^{13,81}

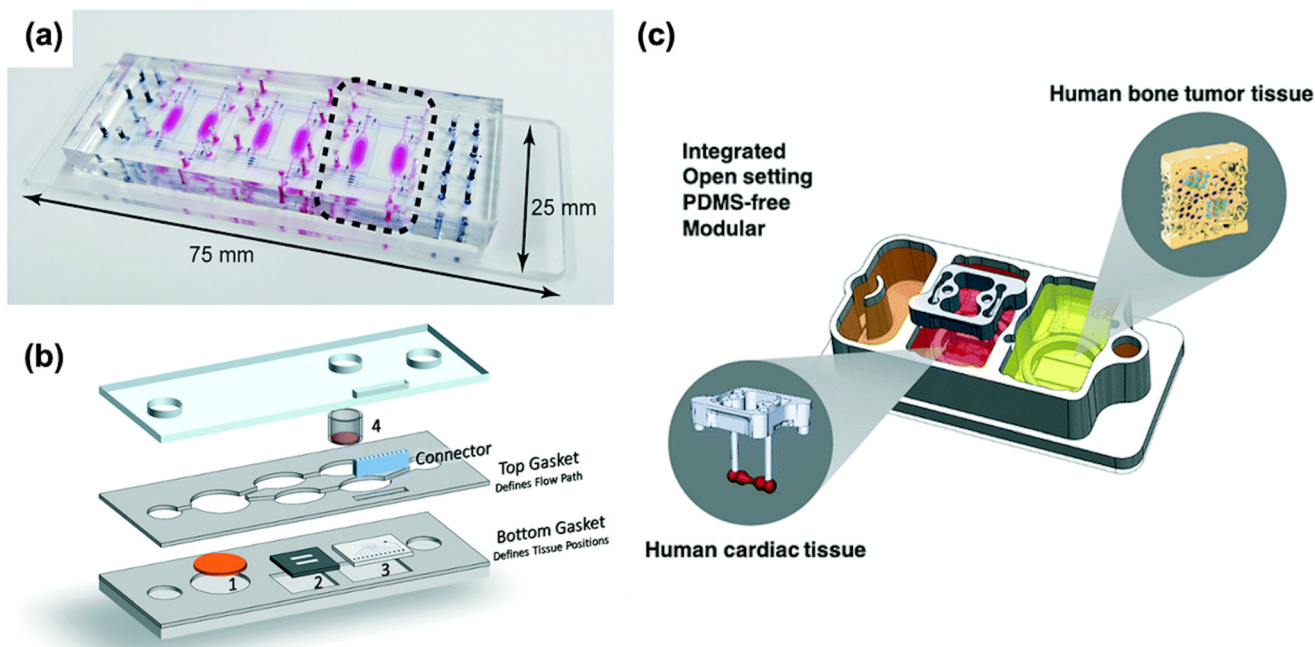


Fig. 6 HoCs as part of multi-OoCs. a) An image of a heart-liver-on-a-chip (HLC) system with three independent co-culture regions on the same chip (reproduced from ref. 431 under Creative Commons 3.0 (CC BY 3.0)). b) A multi-layer HLC comprised of a bottom layer containing the tissues (1 = liver, 2 = cardiomyocytes on a cantilever array, 3 = cardiomyocytes on a microelectrode array, and 4 = skin), middle layer for fluid manipulation, and top layer with access ports (reproduced from ref. 411 with permission from the Royal Society of Chemistry). c) A modular heart-cancer-on-a-chip system with separate chambers for cardiac tissue and Ewing sarcoma (ES) tissue, which can be connected for perfusion studies or kept isolated (reproduced from ref. 360 with permission from the Royal Society of Chem.).

Two of the most common organs affected by drug-induced adverse side effects are the heart and the liver.³⁸⁷ An ingested drug is first absorbed by the small intestine and reaches the liver, where it is metabolized.⁴²⁸ To reach a more comprehensive understanding the pharmacokinetics and pharmacodynamics of a drug and its metabolites, heart–liver-on-a-chip (HLC) systems are being developed (Fig. 6a).^{378,406,414,430–432} Soltantabar *et al.* found doxorubicin treatment to be more damaging to the heart tissue as a result of exposure to not only doxorubicin, but also its metabolite doxorubicinol, which was generated by the hepatocellular carcinoma cells (HepG2) in the chip.^{404,431} One advantage of this HLC platform is the lack of external pumps necessary for fluid perfusion, which reduces the overall device complexity and footprint. However, the use of murine cardiomyocytes (H9c2) and cancerous hepatocytes (HepG2) could lead to different metabolic and pharmacological behavior than that seen with human-derived CMs and normal hepatocytes. To address this discrepancy, Lee-Montiel *et al.* developed an isogenic HLC that utilized hiPSC-CMs and hiPSC-hepatocytes derived from a healthy adult.⁴²⁹ One challenge associated with HLCs is the need for a common culture medium that supports both organ chips. The authors confirmed the liver culture medium to be compatible with the cardiac chip. The multi-organ platform was then used to study drug–drug interaction between cisapride, a gastroprolinctic metabolized by the cytochrome P450 enzyme CYP3A4, and ketoconazole, an antifungal that inhibits CYP3A4. The inherent cardiotoxic effects of cisapride were minimized if cisapride was first passed through the liver tissue before reaching the heart, due to the metabolism of cisapride into non-arrhythmogenic norcisapride. However, the addition of ketoconazole, which inhibits the metabolism of cisapride, resulted in a prolongation of the APD, analogous to clinically observed long QT syndrome. Pires de Mello *et al.* developed a HLC with an added skin surrogate to study common topical pharmaceuticals, including ketoconazole, diclofenac, acetaminophen, and hydrocortisone (Fig. 6b).⁴¹²

The system involved a liver module (containing primary human hepatocytes) and two CM modules (containing hiPSC-CMs), one cantilever array and one microelectrode array for mechanical and electrical characterization, respectively. The skin surrogate was crucial for slowing the delivery of the drug to the main organ chip, which better replicates clinical pharmacokinetics.

Given the well-established cardiotoxicity of many antineoplastics, heart–tumor-on-a-chip systems have been employed to study drug efficacy and cardiotoxicity in one platform. Chramiec *et al.* developed a PDMS-free multi-organ chip to study linsitinib, an experimental anti-cancer drug, for the treatment of Ewing sarcoma (ES), a rare form of bone cancer (Fig. 6c).³⁶⁰ The chip also contained an engineered cardiac tissue chamber to study drug-induced cardiotoxicity in parallel. When tested in the integrated heart–tumor chip, linsitinib (12 $\mu\text{mol L}^{-1}$) was found to have an insignificant effect on cell viability in non-metastatic ES tissue when introduced *via* perfusion, in agreement with observations in clinical trials.

However, direct introduction of the entire volume of linsitinib to the tumor resulted in significant reduction in cell viability, in disagreement with clinical results, which emphasizes the importance of drug perfusion in *in vitro* systems to better replicate clinical scenarios. Furthermore, the perfused experiment showed a slight reduction ($\approx 11\%$) in beat frequency of the cardiac tissue after treatment, which is similar to clinical observations. In another report, Weng *et al.* developed a heart–colon cancer chip with SW620 cells to study doxorubicin and oxaliplatin efficacy and cardiotoxicity.⁴⁰³ Here, the tissues (tumor, endothelial cell, and cardiac) were grown in parallel channels with perpendicular capillaries connecting each channel together, rather than growing each tissue in isolated chambers. One advantage of this configuration is that all three channels can be monitored in the same microscope field of view. Furthermore, the iPSC-EC tissue served as a physiological barrier between the tumor and cardiac tissues. Both DOX and oxaliplatin demonstrated a dose-dependent inhibition of tumor growth in the chip, as well as varying degrees of cardiotoxicity. The IC_{50} of the spontaneous beating rate was measured to be $\approx 0.041 \mu\text{mol L}^{-1}$ for DOX and $33.9 \mu\text{mol L}^{-1}$ for oxaliplatin. Lee *et al.* also tested the efficacy and cardiotoxicity of DOX in a breast cancer (SK-BR-3)-HoC with in-line electrochemical immunosensors to quantify secreted biomarkers.¹⁶² The cardiac tissue was modelled at various stages of fibrosis to replicate breast cancer patients with pre-existing cardiac dysfunctions. DOX treatment was found to decrease beating frequency in both healthy and fibrotic cardiac tissue. Using the electrochemical immunosensors, the healthy cardiac tissue demonstrated a significant increase in troponin T (a marker of myocardial damage) production between 1 and 5 days in culture without DOX, whereas the fibrotic tissue showed no significant change. After treatment with DOX, no significant change in TnT levels were seen between days 1 and 5, while a significant decrease in (human epidermal growth factor receptor 2) HER-2 levels were seen with fibrotic tissue over the same timeline. The authors also compared DOX delivery using magnetic nanoparticles (MNP) with regular free delivery and found MNP delivery to induce less troponin T secretion while maintaining a similar reduction in breast cancer proliferation. Taken together, these examples demonstrate the potential of multi-organ chips to sufficiently model drug metabolism and explore novel therapeutic strategies under controlled and potentially isogenic conditions.

In a paper by McAleer *et al.*, a pumpless liver–heart–tumor-on-a-chip was developed to test both cardiotoxicity and drug efficacy.⁴³² Here, the authors included two cardiac chambers (containing hiPSC-CMs), one with thin film cantilevers and one with a microelectrode array, and two cancer chambers, all downstream of the liver chamber prepared with primary human hepatocytes. The organization of the chambers allows chemotherapeutic drugs to be introduced in a common serum-free medium and metabolized by the liver before reaching the cardiac and cancer chambers, which included breast cancer and multi-drug-resistant vulva cancer cell lines (MCF-7 and SW-962,

respectively). The system showed a reduction in viability of the breast cancer cells after tamoxifen treatment with the liver chamber present, demonstrating the importance of metabolism on drug efficacy. The drug-resistant cancer cells showed a reduction in viability only after treatment with tamoxifen co-administered with verapamil. Off-target effects were observed in the cardiac modules as well, with a reduction in contractile force, conduction velocity, and beat frequency all seen despite without a significant change in cardiomyocyte viability.

6 Standardization in organ-on-a-chip systems

The microphysiological systems (MPS) community have expressed great interest in the development of standards for these platforms. Stemming from the promise of MPS to greatly improve *in vitro* results for toxicity and efficacy testing in drug development, the need for documentary standards (to start with) is of upmost importance in the mind of a number of (bio)pharmaceuticals, and biotechnology companies as well as suppliers of microfluidics and other device components.^{433,434}

The push for standards is illustrated by the efforts of organizations like the Standards Coordinating Body (SCB) in the USA,¹⁴ the North American 3Rs Collaborative,⁴³⁵ the Centre for Alternative to Animal Testing or CAAT (Johns Hopkins University, USA and University of Konstanz, Germany),^{436,437} the Organ-on-a-Chip/Tissue-on-a-Chip Engineering and Efficacy Standardization Working Group (O/TEES WG, USA with collaboration from European Commission),⁴³⁸ CEN-CENELEC (Europe),⁴³⁹ European Organ-on-a-Chip Society (EurOoCS),⁴⁴⁰ and the Microfluidics Association (MFA, International),⁴⁴¹ who have been working to get consensus for specific standard processes, measurements and methods. All these organizations are working with stakeholders from the academia, industry, government research labs and regulatory agencies in the development of guidelines and standards for MPS as well as conferences and publications to further disseminate the work that has been carried out by this community. Although all these efforts are underway, there has always been a concern in certain sectors of the scientific/technology community regarding the possibility of innovation being hindered by the development of standards. However, the development of standards provides the basis for common language and measurement tools that ultimately allow for further advancements to be possible. The areas for standardization discussed below provide a framework for all MPS, but when deciding what specific standards to develop, some of the areas can apply to all MPS and some are more specific. For example, the engineering aspects, combined with protocols, for some of those future standards will likely be applicable to all or almost all MPS. Aspects like flow conditions/control is one that can apply to many MPS uniformly since what could really change from organ to organ is the range at which flow

should be validated. The same way other standards could be applied uniformly, and others will need to be more specific. To make these efforts worthwhile, the community is taking this into consideration and whenever possible standards will be developed in a way that could be applicable as broadly as possible.

The first efforts to develop standards for MPS and related microfluidic devices have so far produced the first three standards in the form of the ISO 22916:2022, "Microfluidic devices – interoperability requirements for dimensions, connections and initial device classification",⁴⁴² the ISO 10991:2023, "Microfluidics – Vocabulary",⁴⁴³ and the ASTM International standard titled "Standard Terminology Relating to Microphysiological Systems", designation F3570-22.¹⁴ However, debates about the next aspects to be standardized are part of the current conversations between and within the different standard developing organizations. Heart MPS is currently being discussed at least by the SCB and O/TEES WG. For each organ model there are numerous areas for the development of standards, and the heart model is no exception. In general, we can think that for all organ models, including Heart MPS, there are three major areas where the development of (documentary) standards will be needed. Those areas are: 1) engineering aspects of microdevices and their components (*i.e.*, the hardware's capabilities and performance); 2) the microenvironment (*i.e.*, the cell adhesive extracellular materials surrounding the cells to provide signals for cell growth, proliferation and more); and 3) protocols to generate functional cells (*e.g.*, mature cardiomyocytes) and their characterization outside and in the microdevice. These are basic areas to establish a baseline of normal cell behavior in the MPS as well as under different perturbations during the assessment of toxicity effects and efficacy of drugs.

In terms of standardization of engineering aspects or hardware capabilities there are several items that apply to Heart MPS as well as to a few other models. For example, heart MPS will require standardized methods to assess the capabilities of integrated components for sensing and actuation. Specifically, it should be considered the development of protocols to reliably measure the mechanical stretching of such components. There should also be considered the development of reliable methods to measure the heart cells beating behavior (*e.g.*, amplitude and frequency). Also, standardization of engineering aspects of the MPS that can modulate the cell state while in the microsystem. One example of engineering aspects to consider is flow control within the system. This parameter is critical since flow affects shear stress, which can exert differences in gene expression in cardiomyocytes, thus producing impaired function and cardiomyopathies.⁴⁴⁴ The microenvironment is an area under a lot of active work to define the optimum ECM and other conditions for cell differentiation and maturation. Components like laminins (*e.g.*, 111, 521 and 511), collagen (*e.g.*, collagen IV), fibronectin and a few other ECM components will need to be part of a combination of

standard materials needed to produce a reliable and reproducible Heart MPS microenvironment.^{444–446} In addition, scaffolds for 3D cultures would need to be considered as part of the microenvironment.⁴⁴⁷ Lastly, protocols to generate functional cells will need to be developed. Those protocols will need to consider the two previous areas discussed above (*i.e.*, engineering aspects and microenvironment) to provide an informed set of procedures for consistently obtaining healthy and mature cardiomyocytes and other cells associated to the organ. Among some of the parameters to be considered are oxygen partial pressure and pH.⁴⁴⁸ Currently, the O/TEES WG is considering cell maturity, beat frequency, conduction velocity, force, and ejection fraction as markers for heart cells behavior and functionality within the MPS. These parameters refer to protocols to assess functional cells, mainly physical measurements of cell processes. However, other efforts would be expected to take a different route and investigate other parameters such as ATP content, cardiac troponin and chemical markers for cardiac cells function (*i.e.*, protocols focusing on biochemical assessment).

7 Conclusions and outlook

The heart is responsible for efficiently delivering blood, nutrients, and oxygen to tissue and organs throughout the body. Given its outsize role in regulating physiological homeostasis, there is great interest in holistically understanding cardiovascular physiology and pathophysiology. Historically, improving this understanding has been challenging because of a lack of sufficiently relevant human models. HoCs systems have advanced tremendously over the past decade, coinciding with improvements to tissue engineering, gene editing, and especially the advent of human-induced pluripotent stem cells. These advancements have brought about new applications for HoCs as discussed in this review, and with that, new companies working to commercialize this technology. Nonetheless, significant work remains to be done before HoCs, and OoCs more broadly, become commonly used in pre-clinical testing.

One substantial hurdle that needs to be addressed is the challenge of tissue immaturity in these systems. This is especially a problem for stem-cell derived cardiac cells, which typically express a fetal-like phenotype. As discussed in section 4.1.1, numerous methods for improving maturity have been reported, including mechanical conditions, electrical stimulation, biochemical treatment, and cell co-culture. As a result, more human-relevant assessments of disease progression and drug toxicity have been made. We expect to see continued work in the area of tissue maturation as HoCs advance along the technological development pathway. Despite advances in tissue maturation, the ability to holistically model the comorbidities that often accompany cardiovascular dysfunction in adults is still in its infancy. Another area that remains a bottleneck to widespread commercialization is chip material. PDMS is the most widely

used material for fabricating microfluidic chips for OoCs. Although PDMS has advantages in terms of gas permeability, biocompatibility, optical transparency, and ease of patterning, it suffers from a number of key drawbacks that could ultimately limit its use as OoC technology progresses. Without any form of automation, the assembly of PDMS OoCs can be tedious and artisanal. Another key issue particularly relevant for drug development is the absorption of many small molecules in PDMS, which can skew toxicity and efficacy results if not addressed and/or accounted for.⁸⁸ Furthermore, PDMS is relatively expensive when compared to other materials currently used in cell biology, such as polystyrene, poly(methyl methacrylate), and other polymers. These materials are more conducive to industrial-scale manufacturing processes, such as 3D printing and injection molding. Until OoCs can be quickly and inexpensively prepared using large-scale manufacturing methods, commercialization of the technology will likely be limited.

We believe that standardization will play a critical role in many of the areas associated with OoCs, making widespread adoption of OoCs more appealing. From a production and design perspective, standardized materials, geometries, and treatments will enable easier cross-platform comparison. Cell biology has benefited tremendously from the geometric standardization of tissue culture vessels, allowing for more universal protocols and viable high-throughput instrumentation. Reporting of a common set of metrics will also accelerate progress and make commercialization easier. By establishing a common set of metrics, not only can researchers compare their work to that of others, but also approval from regulatory agencies should become less cumbersome as a result. Rigorous methods for benchmarking results from OoCs with animal studies will most likely lead to faster and broader acceptance of a given OoC model as a comparable, if not better, alternative to the equivalent animal model.

Regulatory agencies, such as the US Food and Drug Administration (FDA), have already lifted the requirement of animal testing when seeking approval of a new drug, which opens the door for new and novel alternatives, including OoCs, to replace them. As a result, one application in which OoCs could find widespread acceptance is in the re-purposing or accelerated approval of drugs and therapeutics during public health emergencies, where animal studies and clinical trials may be too time-consuming. The COVID-19 pandemic is a recent example where numerous existing drugs, such as hydroxychloroquine and azithromycin, were claimed to treat or manage COVID-19. However, these claims were ultimately nullified after users experienced a number of adverse side effects, including kidney injury and liver damage.⁴⁴⁹ While these instances are not entirely avoidable, OoCs could help improve and accelerate drug re-purposing and dissemination of information during rapidly changing situations, such as a global pandemic or other public health emergency.

Finally, we expect to see the incorporation of on-board analytical tools increase as OoCs become more sophisticated.

Biosensors, including electrochemical and optical, will become more heavily integrated in OoC systems as real-time understanding of tissue behavior is needed and traditional end-point assays lack the ability to provide the necessary information. Real-time measurements are especially important for dynamic samples, such as cardiac tissue, which can change behavior (e.g., beating frequency) in a matter of seconds. Moreover, integration of auxiliary sensors, such as temperature and pH, among others, will assist in standardization and improve data robustness. As costs decrease, and sensor integration and on-board analysis improves, OoC systems could see more decentralized usage outside of a clinical environment in more resource-limited settings.

We envision marked advancements to OoCs in the near future, driven by increased human relevance, decreased costs, and robust standardization. As tissue engineering, scalable manufacturing, gene editing and analytical methods improve, more complex models can be realized leading to a deeper and more personalized mechanistic insight of cardiophysiology, disease, and toxicity. As models improve, we expect better agreement between results obtained from OoCs and clinical observations, leading to more widespread adoption of OoC models as a suitable replacement for animal models. While widespread adoption and commercialization will take time, it is likely that OoCs will find use in certain applications sooner than others. For example, rapid drug screening during public health emergencies or rare disease treatment where there may otherwise be difficult in establishing a trial cohort are just some of the applications that could benefit from prompt adoption of OoCs. Overall, OoCs are only in their infancy and are expected to cause a paradigm shift in the way pre-clinical and clinical assessments of disease treatment and drug efficacy and toxicity are carried out.

Author contributions

D. B. and D. R. R. both contributed to literature review, writing, and editing.

Conflicts of interest

There are no conflicts to declare.

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