

1 **Title: Electronic blood vessel**

2

3 **Authors:** Shiyu Cheng^{1,2,4,5}, Chen Hang^{1,2,5}, Li Ding^{1,2,3,5}, Liujun Jia³, Lixue Tang²,

4 Lei Mou^{1,2}, Jie Qi^{1,2}, Ruihua Dong^{1,2}, Wenfu Zheng², Yan Zhang^{3*}, Xingyu Jiang^{1,2,6*}

5 **Affiliations:**

6 ¹Department of Biomedical Engineering and Shenzhen Bay Laboratory, Southern

7 University of Science and Technology, Shenzhen, Guangdong 518055, China.

8 ²National Center for NanoScience and Technology, University of Chinese Academy
9 of Sciences, Beijing 100190, China.

10 ³State Key Laboratory of Cardiovascular Disease, Fuwai Hospital, National Center
11 for Cardiovascular Diseases, Chinese Academy of Medical Sciences and Peking
12 Union Medical College, Beijing 100037, China.

13 ⁴Current address: Institute of Bioengineering, School of Engineering, Ecole
14 Polytechnique Federale de Lausanne, Lausanne 1015, Switzerland.

15 ⁵These authors contributed equally.

16 ⁶Lead Contact

17 *Correspondence: jiang@sustech.edu.cn; waikezhangyan@126.com

18

19 **Summary**

20 Advances in bioelectronics have great potential to address unsolvable biomedical
21 problems in the cardiovascular system. By using poly(L-lactide-co-ε-caprolactone)
22 (PLC) that encapsulates the liquid metal to make flexible and bio-degradable
23 electrical circuitry, we develop an electronic blood vessel that can integrate flexible
24 electronics with three layers of blood vessel cells, to mimic and go beyond the natural
25 blood vessel. It can improve the endothelization process through electrical stimulation
26 and can enable controlled gene delivery into specific part of the blood vessel via
27 electroporation. The electronic blood vessel has excellent biocompatibility in the
28 vascular system and shows great patency three months post-implantation in a rabbit
29 model. The electronic blood vessel would be an ideal platform to enable diagnostics
30 and treatments in the cardiovascular system and can greatly empower personalized
31 medicine by creating a direct link of vascular tissue-machine interface.

32 **Keywords:** Metal polymer conductor, blood vessel, electrical stimulation,
33 electroporation, bioelectronics, vascular interface.

34

35 **Introduction**

36 Cardiovascular diseases remain the number one cause of mortality worldwide¹. In
37 treating cardiovascular diseases via coronary artery bypass grafting (CABG) surgery,
38 none existing small diameter (<6 mm) tissue engineered blood vessel (TEBV) has met
39 clinical demands². To fabricate the TEBV, a range of approaches, such as
40 decellularized matrix³⁻⁷, self-assembly cell sheets⁸⁻¹¹, bioactive and biomimetic
41 materials¹²⁻¹⁴, were developed and clinically investigated in recent years. However,
42 most of these methods only serve as scaffolds to provide mechanical support and
43 mainly rely on the remodeling process by the host tissue and present significant
44 limitations in helping the regeneration of neo blood vessel. Thus far, none of them
45 have achieved satisfactory clinical results. Specifically, a complex interplay between
46 the blood flow and the TEBV can often cause inflammatory responses, resulting in
47 thrombosis, neointimal hyperplasia or smooth muscle cell accumulation nearby the
48 scaffold^{2,15}, in different pathological stages. To address these issues, next-generation
49 TEBV should not only function as scaffolds to provide the mechanical support and
50 facilitate host cell recruitments, but also have the capability to actively respond to and
51 couple with the native remodeling process in order to provide adaptive treatments
52 after implantation.

53 Combining living tissues with flexible electronics^{16,17} could endow the conventional
54 TEBV more functionalities and capabilities to overcome existing biomedical
55 problems, such as precision diagnostics, by *in situ* sensing the blood flow and
56 temperature, and treatments by therapeutic drug or gene delivery^{18,19}. In previous

57 works, we have developed many approaches to fabricate structures that mimic natural
58 blood vessel with multiple types of blood vessel cells in different layers, including
59 stress-induced self-rolling membrane^{13,20-22} and layer by layer technique²³. Recently,
60 we have developed a printable metal-polymer conductor (MPC) which exhibited
61 excellent compatibility and high stretchability^{24,25}. Here, we develop an electronic
62 blood vessel that integrates flexible electrodes into the bio-degradable scaffold by
63 combining the liquid metal with poly(L-lactide-co-ε-caprolactone) (PLC) into a
64 metal-polymer conductor. As a proof of concept, we used the electronic blood vessel
65 to carry out *in vitro* electrical stimulation and electroporation. By electrical
66 stimulation, the electronic blood vessel can effectively promote cell proliferation and
67 migration in a wound healing model. It can also *in situ* deliver the GFP DNA plasmid
68 into three kinds of blood vessel cells via electroporation. We evaluated the efficacy
69 and biosafety of the electronic blood vessel in the vascular system through a three-
70 month *in vivo* study by using a rabbit carotid artery replacement model and confirmed
71 its patency by using ultrasound imaging and arteriography. Our results pave the way
72 to integrating flexible, degradable bio-electronics into the vascular system, which can
73 serve as a platform to carry out further treatments, such as gene therapies, electrical
74 stimulation, electronically controlled drug releases.

75

76 **Results**

77 **Fabrication of the electronic blood vessel.**

78 We fabricated the electronic blood vessel (**Figure 1A**) by rolling up a PLC based
79 metal-polymer conductor (MPC-PLC) membrane (**Figure 1B**) with the assistance of a
80 polytetrafluoroethylene (PTFE) mandrel. The MPC circuit was well distributed in the
81 three dimensional (3D) multilayered tubular structure. The inner diameter of the
82 electronic blood vessel was around 2 mm (**Figure 2A**) and the minimum diameter
83 could be around 0.5 mm (**Figure S1A**). The MPC-PLC membrane is flexible and
84 degradable and the MPC circuit is conductive (**Figure 1B-D**). The conductivity of the
85 MPC circuit is about 8×10^3 S cm⁻¹ and the $\Delta R/R$ of the circuit remained constant after
86 around 1000 cycles of bending and rubbing (**Figure 1C**). The PLC is projected to be
87 entirely degraded around 1~2 years by the manufacturer. The MPC-PLC membrane
88 underwent a mass loss of around 10% during 8-week incubation in the phosphate-
89 buffered saline (PBS, 37 °C) (**Figure 1D**). We observed a relatively quick degradation
90 in the first week (**Figure S2**). To fabricate the MPC-PLC membrane, we screen-
91 printed the conductive ink on the polyethylene terephthalate (PET) membrane (**Figure**
92 **1E**). The electrode design was optimized for electroporation and electrical
93 stimulation. By preparing different electrode designs, we could either target individual
94 blood vessel layers (tunica intima/media/adventitia) with the electrodes distributed in
95 specific areas or target all three layers with the full electrode (**Figure S1B-C**). We
96 prepared the liquid metal conductive ink by sonicating the mixture of Gallium-Indium
97 alloy (EGAln, ≥99.99%, Sigma, US) and the volatile solvent (1-decanol, Macklin,

98 Shanghai) (**Figure 1F, H**). The liquid metal particles (LMPs) exhibit a core-shell
99 structure, where the core is the Ga-In alloy and the shell is the Ga-In Oxide (**Figure**
100 **1G**). The diameter of the LMPs is around 2 μm (**Figure 1J**). We embedded the LMPs-
101 based circuit with the PLC solution (5 wt % in CH_2Cl_2) and peeled the MPC-PLC
102 membrane off the PET substrate after evaporation of the solvent in a chemical hood
103 (**Figure 1E**). The thickness of the MPC-PLC membrane is about 50 μm and it is
104 tunable by changing the volume of the PLC solution. The LMPs could be broken
105 during the process of peeling and release the Ga-In alloy to make the circuit
106 conductive²⁵. We confirmed the structure of the conductive circuit via corrosion of the
107 Ga-In alloy by adding excessive hydrochloric acid (HCl, Beijing Chemical Works,
108 China). The LMPs were evenly distributed in the cellular PLC host (**Figure 1K, L**).
109 The liquid metal could form a coherent conductive pathway in the polymeric host.
110

111 ***In vitro* characterization of the electronic blood vessel.**

112 To evaluate biocompatibility of the electronic blood vessel, we used the microfluidic
113 technology to realize accurate 3D pattern of three kinds of blood vessel cells in a
114 natural blood vessel mimicking fashion. By employing a multichannel microfluidic
115 chip, we delivered human umbilical vein endothelial cells (HUVECs, blue), human
116 aortic smooth muscle cells (SMCs, green), human aortic fibroblasts (HAFs, red)
117 sequentially on the MPC-PLC membrane (**Figure 2A-E**). We designed the width of
118 each channel to match the circumference of each layer of the tube based on thickness
119 of the MPC-PLC membrane and the diameter of the tube (**Figure 2B**). To distinguish
120 different cell types, we stained the HUVECs, SMCs, HAFs with different fluorescent
121 dyes (HUVECs: Cell Tracker Violet; SMCs: Cell Tracker Green; HAFs: Cell Tracker
122 Deep red) before seeding them into the microfluidic chip. After 1 d incubation in
123 culture medium (Dulbecco's modified eagle medium, DMEM, 10% FBS, 37 °C, 5%
124 CO₂), cells attached on the MPC-PLC membrane (**Figure 2D, H**). We peeled the
125 microfluidic chip off the cell-laden MPC-PLC membrane and rolled it up with a
126 PTFE mandrel, forming a 3D multi-layered tubular structure (**Figure 2E-G**) with
127 HUVECs, SMCs, and HAFs evenly distributed in the inner layer, middle layer, and
128 outer layer, respectively (**Figure 2I, J**). This structure well mimics the structure of the
129 natural blood vessel. To better understand the blood vessel cells distributed in
130 different layers of the electronic blood vessel, we stained HUVECs, SMCs, HAFs,
131 and the MPC-PLC layer with CellTracker DiO, CellTracker DiI, CellTracker DiD,
132 and CellTracker Blue to show the relative distribution of different layers (**Figure S3**).

133 We used a biomedical fibrin glue to facilitate the combination of different layers¹⁸. We
134 incubated the cell-laden electronic blood vessel for 14 d and stained the cells with the
135 Calcein AM green. The evenly distributed green color on the MPC-PLC membrane
136 indicated that the cells exhibited high viability after a 14 d culture (**Figure 2K**). We
137 measured the transport of ions through the MPC-PLC membrane to further evaluate
138 and quantify the permeability of the electronic blood vessel (**Figure S4**). Ca^{2+} , Fe^{3+} ,
139 Mg^{2+} could permeate the electronic blood vessel over time. We also conducted a
140 hemolysis test, which showed that the electronic blood vessel exhibited very good
141 blood biocompatibility (**Figure 2L**). The *in vitro* characterization demonstrated that
142 the electronic blood vessel exhibited excellent biocompatibility and we then
143 conducted further functional tests of the embedded MPC circuits, i.e. electrical
144 stimulation and electroporation.

145
146 ***In vitro* electrical stimulation promotes HUVEC proliferation and migration**
147 To prove the functionality of the electronic blood vessel, we carried out *in vitro*
148 electrical stimulation to improve proliferation and migration of HUVECs. Direct
149 current (DC) electric field has been shown to effectively increase the angiogenesis *in*
150 *vitro* and *in vivo*²⁶. We patterned HUVECs on the MPC-PLC membrane (DMEM,
151 10% FBS, 37 °C, 5% CO_2) by using a multi-chamber PDMS chip. The initial number
152 of cells in each chamber was the same (3×10^4). After 12 h incubation, we applied
153 different DC voltages to yield different electrical field strength: 25, 50, 75 100, 200,
154 and 400 mV mm⁻¹, respectively (**Figure 3A, B**). After 2 d incubation and electrical

155 stimulation, we randomly selected 6 different domains on each sample and analyzed
156 by using a laser scanning confocal microscopy (LSCM, LSM 710, Zeiss, Germany).
157 We stained nuclei (blue) with Hoechst 33342 (Invitrogen, USA) and stained the living
158 and dead cells with Calcein-AM (green) and PI (red), respectively. The green cells
159 occupied the whole domain, indicating that electrical stimulation did not hurt
160 proliferation of HUVECs (**Figure 3C**). We counted the number of nuclei by using
161 ImageJ. The cell number under 50 mV mm^{-1} was highest, about 2.4 times that of the
162 control (**Figure 3E**). We used the CCK-8 kit to confirm this conclusion (**Figure 3F**).
163 We speculated that the DC electric field had selectively regulated production of
164 certain growth factors and cytokines important for angiogenesis.

165 We explored the migration of HUVECs under different DC electric field strengths.
166 We made a scratch on a PDMS substrate by using a $10 \mu\text{l}$ tip. After application of an
167 electric field of 50 mV mm^{-1} , HUVECs migrated $750 \mu\text{m}$ and the wound completely
168 healed after 24 h (**Figure 3D**). Different strengths had enhanced migration differently
169 compared to the control group without electrical stimulation (**Figure 3G**). The *in vitro*
170 DC electrical stimulation thus proved to have effectively promoted the proliferation
171 and migration of HUVECs.

172 We further evaluated the effectiveness of electrical stimulation in a 3D model of
173 endothelialization. We patterned the HUVECs on the MPC-PLC membrane and made a
174 scratch on the cell-laden membrane by using a $10 \mu\text{l}$ tip. We transformed the 2D cell-
175 laden membrane into 3D cell-laden structure and connected it to the electrochemical
176 station to test the proliferation and migration. We applied an electric field of 50 mV

177 mm⁻¹ on multiple samples and we observed the proliferation and migration at different
178 time points. We stained the living and dead cells with Calcein-AM (green) and PI
179 (red). We counted the cell number and the cell density was higher than the control
180 group (Figure 3H). The HUVECs formed a complete endothelial layer after 24 h
181 (Figure 3I). To better evaluate the biocompatibility of the MPC circuit under electrical
182 stimulation, we extended the electrical stimulation time to 10 days and the live/dead
183 staining showed that cells exhibited excellent viability (Figure S5).

184

185 ***In vitro* electroporation**

186 To further prove the functionality of the electronic blood vessel, we designed multiple
187 circuit patterns for electroporation, being able to target different pathological issues in
188 different layers of blood vessel cells (**Figure S1B**). We conducted the electroporation
189 with a circuit pattern that could target all three layers. We seeded the cells onto the
190 MPC-PLC membrane and transformed it into 3D tubular structure for electroporation.
191 We immersed the 3D cell-laden electronic blood vessel in the GFP plasmid DNA
192 solution for 10 min before electroporation. The GFP plasmid DNA could also be
193 lyophilized onto the MPC-PLC membrane before seeding cells (**Figure S7A**) and
194 transforming to 3D structure for electroporation. We connected the 3D cell-laden
195 electronic blood vessel with an electroporator (BTX, CM630, US) to generate DC
196 pulses (**Figure 4A**) and achieved the delivery of the green fluorescent protein (GFP)
197 DNA plasmid in three kinds of blood vessel cells (**Figure 4B**). To optimize the
198 performance of the electronic blood vessel, we found two major parameters

199 determining the efficacy of electroporation, including voltage and pulse duration. We
200 conducted the electroporation with different voltages (40 V/60 V/80 V) and pulse
201 durations (100 μ s/1 ms). If the voltage was too low, it would cause low efficacy or no
202 transfection; if the voltage was too high or if the pulse duration was too long, it would
203 cause low efficacy and cell death (**Figure S6**). To realize the optimal efficacy, we
204 exerted a square wave with the voltage of 60 V, pulse duration of 100 μ s, and pulse
205 interval of 1 s for 5 times. We delivered GFP plasmid DNA into three kinds of blood
206 vessel cells and the GFP DNA realized expression with more than 95% of cells
207 showing green fluorescence (**Figure 4B**). We observed successful expression of GFP
208 among all three layers of the blood vessel cells and exhibited a uniform 3D
209 distribution in the electronic blood vessel (**Figure 4C**). To evaluate the potential of
210 the electronic blood vessel for *in vivo* electroporation, we lyophilized the GFP
211 plasmids DNA on the MPC-PLC membrane to test its effectiveness (**Figure S7A**). We
212 carried out electroporation by attaching the plasmids-laden MPC-PLC membrane to
213 an isolated rabbit vascular tissue with the voltage of 60 V, pulse duration of 100 μ s,
214 and pulse interval of 1 s for 5 times. We observed successful expression of GFP in the
215 isolated rabbit vascular tissue (**Figure S7A**) after 2 d incubation. These promising *in*
216 *vitro* results encouraged us to carry out the *in vivo* tests of the electronic blood vessel.
217

218 **Mechanical properties of the electronic blood vessel**

219 To find out whether the electronic blood vessel is suitable for *in vivo* implantation, we
220 measured the mechanical properties, including stress-strain curve, compliance, and

221 burst pressure, of the electronic blood vessel with a diameter of 2 mm prior to
222 implantation (**Figure 5A-F and Figure S8**). The elastic modulus of the electronic
223 blood vessel is about 130 MPa, the value is much higher than the native carotid artery.
224 The ultimate tensile strength of the electronic blood vessel is about 27 MPa, the value
225 is also much higher than that of the native carotid artery. The initial compliance of the
226 electronic blood vessel (n= 5) is about 5% per 100 mmHg in the range of 80-120
227 mmHg, the value is apparently below that of the native carotid artery (n= 3). The
228 burst pressure of the electronic blood vessel (n= 5) is about 2800 mmHg, the value is
229 similar to that of the native carotid artery (n= 3). The elongation at break of the
230 electronic blood vessel is about 650% (n= 5), which is the twice as the value of native
231 carotid artery (n= 3). The mechanical property of electronic blood vessel was
232 considered robust enough for implantation.

233

234 ***In vivo* implantation in rabbits and *in situ* monitoring**

235 To investigate the electronic blood vessel as a vascular implant, we chose the New
236 Zealand rabbit (age: 200-300 days, body weight: 3-4 kg) as the animal model and
237 replaced the native carotid artery with the electronic blood vessel (**Figure 6A-C**). To
238 avoid possible immunological response of the host tissue, we used the acellular
239 electronic blood vessel in the preliminary *in vivo* study. We *in situ* monitored the
240 implanted electronic blood vessel by doppler ultrasound imaging (**Figure 6D-I**) and
241 the arteriography (**Figure 6J, K**). Doppler ultrasound imaging showed that the
242 electronic blood vessel allowed for good blood flow 3 months post-implantation

243 (Figure 6D-G and Video S1). The asymmetric velocity curve synchronized with the
244 ultrasonic pulses indicated that the signal is from the carotid artery rather than the
245 vein (Figure 6D). The diameter of the electronic blood vessel remained at a relatively
246 constant value, about 2.3 mm, during the half month to three months post-
247 implantation (Figure 6G, H). The mean velocity of the blood flow in different
248 samples at different time point was about 0.47 m s^{-1} , which was in the range of the
249 normal value (Figure 6I). As the gold standard of the blood vessel patency,
250 arteriography showed that the electronic blood vessel matched with the native carotid
251 artery very well and allowed for excellent blood flow (Figure 6J, K, Figure S9, and
252 Video S2). There was no sign of narrowing. The electronic blood vessel allows
253 straightforward visualization under arteriography because the liquid metal-based
254 circuitry has sufficiently high contrast over host tissues (Figure 6J). The red frame
255 outlined the electronic blood vessel with an alternate strip structure from the MPC
256 circuit.

257

258 ***Ex vivo* study**

259 We dissected all the implanted electronic blood vessels three months post-
260 implantation for characterization. The lumen and the outer surfaces of the explanted
261 electronic blood vessel were smooth, covered by the remodeling tissues (Figure 7A,
262 B). The diameter of the native blood vessel significantly reduced due to the lack of the
263 blood pressure, whereas the electronic blood vessel remained the same as before
264 (Figure 7C). We observed the micro structure of the circuit by using the SEM after

265 explanting the electronic blood vessel from rabbit. The MPC-PLC membrane still
266 maintained interdigitated structure with the MPC circuit and PLC host (**Figure 7E**).
267 There was a layer of neo-tissue formed, which well covered the MPC-PLC membrane
268 (**Figure 7F-H**). We could also see some red blood cells on the top of the circuits,
269 which were similar in number to those on native blood vessels (**Figure 7D**). We tested
270 the conductivity of the circuit in the electronic blood vessel. The MPC circuit was still
271 conductive and the conductivity was around $7.2 * 10^3$ S cm⁻¹.

272 To study the histological changes of the implanted electronic blood vessels, we
273 performed histological staining of the electronic blood vessels, setting native carotid
274 blood vessel as a positive control. Hematoxylin/eosin (H&E) staining (**Figure 7I**) of
275 the cross section of the electronic blood vessel is round-shaped, continuous, red,
276 which is similar as the native blood vessel. The three-layered MPC-PLC membrane
277 merged into one intact layer with secretion of substantial extracellular matrix between
278 the different layers of electronic blood vessel. We could clearly see dark blue nuclei
279 (red arrows in Figure 7I) in all the layers, which indicated successful migration and
280 infiltration of host cells into the electronic blood vessel. We compared the cell density
281 in the electronic blood vessel with the native carotid. The cell density of the electronic
282 blood vessel was around 400 cells mm⁻² whereas that of the native carotid was around
283 535 cells mm⁻². More importantly, a dense layer of cells with curved structure were
284 formed in the lumen of the electronic blood vessel, which indicated the excellent
285 endothelialization and thus ensured good blood flow. To further confirm the components
286 in the implanted electronic blood vessel, we performed Masson's trichrome (**Figure**

287 **7J**) and Verhoeff's staining (**Figure 7K**). Masson's trichrome staining can stain and
288 assess keratin and muscle fibers (red) or collagen (blue) and Verhoeff's staining can
289 stain and assess the presence of elastin fibers. Masson's trichrome and Verhoeff's
290 staining showed well-distributed collagen and elastin fibers both inside the layers and
291 between different layers, indicating appropriate remodeling. Compared to the
292 abundance level of extracellular matrix in the native blood vessel, the electronic blood
293 vessel still requires more time for material degradation and tissue remodeling. These
294 results indicated that the electronic blood vessel might still be functioning with the
295 presence of conductive materials and host remodeling 3 months post-implantation.

296 To investigate the influence of the electronic blood vessel on the host, we
297 performed the cross section and histological staining of the major organs, including
298 heart, liver, spleen, lung, and kidney, together with the dissection of the implanted
299 electronic blood vessel 3 months post-implantation. The H&E staining and Masson's
300 Chrome staining showed that there was no significant pathological changes or
301 inflammatory responses in these organs (**Figure S10**). To evaluate whether there were
302 chronic inflammation or infection, we conducted the ELISA assay of three important
303 proteins in the blood, including interleukin-6 (IL-6), procalcitonin (PCT), and C-
304 reactive protein (CRP) (**Figure S11**). The concentrations of IL-6 and PCT were not
305 higher than the normal value of healthy rabbits (red dotted lines). The concentration
306 of CRP was higher than the normal value of healthy rabbits (red dotted line). All three
307 indexes decreased over time, a tendency that was expected. The results showed that
308 most of the indexes were in the normal range and there were no significant

309 pathological changes or inflammatory responses. We also conducted the complete
310 blood count of the rabbit, including white blood cell count, absolute neutrophil count,
311 and absolute lymphocyte count, most of which were in the normal range over time
312 (**Figure S12**). These results confirmed that as an implant for vascular system, the
313 electronic blood vessel had no significant detriment to the host.

314

315 **Discussion**

316 None of the existing small-diameter TEBVs has met the demands in treating
317 cardiovascular diseases. Conventional TEBVs can be greatly improved to provide
318 next-generation treatment by integrating with flexible bioelectronics. In this work, we
319 report an electronic blood vessel with excellent biocompatibility, flexibility,
320 mechanical strength, and degradability by combining the MPC with a US FDA-
321 approved biodegradable polymer. As a proof of concept, we verified that the
322 electronic blood vessel can accelerate the HUVEC proliferation and migration by
323 electrical stimulation, thus facilitating the endothelialization process, which is important
324 to an engineered vascular conduit in preventing early thrombosis. Because most
325 TEBVs occluded with 2 weeks post-implantation, our electronic blood vessel, with a
326 patency of at least 12 weeks, is of great promise for clinical application. We also
327 showcased that it could be used to perform *in situ* gene delivery via electroporation,
328 which laid a foundation for future design and optimization such that we can carry out
329 further gene therapies targeting different pathological problems after implantation.

330 The electronic blood vessel has a high level of safety. The liquid metal^{16,27} has been
331 proven to be highly biocompatible and the PLC has been approved by the US FDA for
332 implants²⁸. We validated its biosafety in the vascular system by a 3-month
333 implantation in a rabbit carotid artery model. Both the *in situ* monitoring, including
334 doppler ultrasound imaging and arteriography, and *ex vivo* study demonstrated that it
335 was safe as an electronic implant in the vascular system and at the body level. The
336 electronic blood vessel exhibited higher strength than the native blood vessel. The

337 potential harm of a rigid synthetic blood vessel is the mismatch with host tissue after
338 implantation. However, from the *in vivo* results, these discrepancies did not bring
339 about any major issues and the electronic blood vessel matched very well with the
340 host carotid artery during the time of *in situ* monitoring (Figure 6J, K, Video S2). The
341 reasons why we chose liquid metal in the electronic blood vessel are as following: i)
342 comparing with gold or platinum, the Ga-In liquid metal allows superior flexibility
343 and stretchability while maintains good conductivity, which is critical for an artificial
344 blood vessel to adapt to the deformation due to the rhythmic beating; ii) it exhibits
345 excellent cytocompatibility and blood compatibility according to our results; iii)
346 compared to other sophisticated microfabrication techniques, using the screen printing
347 technique is much more straightforward and could enable industrial-scale mass
348 production in a cost-effective manner.

349 As a vascular substitute, the electronic blood vessel breaks through the limitations
350 of the existing vascular scaffold by endowing the electrical function to conventional
351 biodegradable TEBV and provides us a new platform for tackling the problems
352 threatening the small-diameter blood vessel. By integrating with other electronic
353 devices, the electronic blood vessel can provide various treatments, such as electrical
354 stimulation, electroporation, electrically controlled drug release, and so forth. When
355 combining with emerging technologies such as artificial intelligence, it can greatly
356 boost future personalized medicine by bridging the vascular tissue-machine interface
357 and empowering the health data collection and storage, such as blood velocity, blood
358 pressure, and blood glucose level. In the future, optimizing its function and creating a

359 multifunctional electronic blood vessel can greatly benefit human cardiovascular
360 health.

361 **Conclusions**

362 In this work, by integrating the liquid metal-based conducting circuitry with a
363 biodegradable polymer, we develop an electronic blood vessel, with excellent
364 biocompatibility, flexibility, conductivity, mechanical strength, and degradability, that
365 enables *in situ* electrical stimulation to facilitate the endothelization process and
366 electroporation to deliver genes in specific layers of blood vessel cells. It exhibited
367 excellent patency and biosafety 3 months post-implantation in the vascular system of
368 a rabbit model. In the future, the electronic blood vessel can be integrated with other
369 electronic components and devices to enable diagnostic and therapeutic functions and
370 greatly empower personalized medicine by creating a direct link of vascular tissue-
371 machine interface.

372

373 **Supplemental Information**

374 Supplemental Information can be found online.

375 **Acknowledgements**

376 This study was supported by the National Key R&D Program of China

377 (2018YFA0902600, 2017YFA0205901), the National Natural Science Foundation of

378 China (21535001, 81730051, 21761142006, 51973045), the Chinese Academy of

379 Sciences (QYZDJ-SSW-SLH039, 121D11KYSB20170026, XDA16020902),

380 Shenzhen Bay Laboratory (SZBL2019062801004), the Tencent Foundation through

381 the XPLORER PRIZE, Beijing Science and Technology Plan Project

382 (Z191100007619053), Postgraduate Innovation Foundation of Peking Union Medical

383 College (2019-1002-28) and Teaching Reform Foundation of Postgraduate Education

384 in Peking Union Medical College (10023201900202). We thank Dr. Zewen Wei and

385 Dr. Deyao Zhao for discussions on the electroporation experiment. We thank Mrs.

386 Barbara Althaus at EPFL for her advice in writing the manuscript.

387 **Author Contributions**

388 X. J. and S. C. conceived the idea. S. C. designed the experiments, fabricated the

389 blood vessel, conducted the characterization, *in vitro* tests, mechanical tests, *in vivo*

390 tests and *ex vivo* tests, analyzed the data, and wrote the manuscript. C. H. fabricated

391 the blood vessel, performed *in vitro* tests, mechanical tests, and *in vivo* tests. L. D.

392 performed the *in vivo* and *ex vivo* tests. S. C., C. H., and L. D. contributed equally. L.

393 T. assisted with the preparation of the liquid metal particles, MPC-PLC membrane

394 and the electroporation experiments. L. J. and Y. Z. performed the implantation of the

395 electronic blood vessel, doppler ultrasound imaging and arteriography. L. M.
396 conducted the ELISA assay. J. Q. and R. D. assisted the TEM characterization. W. Z.
397 gave inputs on the *in vitro* tests and *ex vivo* tests. Y. Z. supervised the *in vivo* and *ex*
398 *vivo* study. X. J. supervised the project, and revised the manuscript. All the authors
399 took part in the discussion and writing.

400 **Declaration of Interests**

401 S. C. and X. J. declare financial interest in form of a patent application. Other authors
402 declare no competing financial interests.

403 **Data Availability**

404 All data are available from the corresponding authors upon reasonable request.

405

406 **References and Notes**

407 1. WHO | World Health Statistics 2019: Monitoring health for the SDGs. *World
408 Heal. Organ.* (2019).

409 2. Seifu, D. G., Purnama, A., Mequanint, K. & Mantovani, D. (2013). Small-
410 diameter vascular tissue engineering. *Nat. Rev. Cardiol.* **10**, 410–421.

411 3. Quint, C. *et al.* (2011). Decellularized tissue-engineered blood vessel as an
412 arterial conduit. *Proc. Natl. Acad. Sci.* **108**, 9214–9219.

413 4. Dahl, S. L. M. *et al.* (2011). Readily available tissue-engineered vascular
414 grafts. *Sci. Transl. Med.* **3**, 68ra9.

415 5. Lawson, J. H. *et al.* (2016). Bioengineered human acellular vessels for dialysis
416 access in patients with end-stage renal disease: two phase 2 single-arm trials.
417 *Lancet* **387**, 2026–2034.

418 6. Zhu, M. *et al.* (2019). In vivo engineered extracellular matrix scaffolds with
419 instructive niches for oriented tissue regeneration. *Nat. Commun.* **10**, 4620.

420 7. Kirkton, R. D. *et al.* (2019). Bioengineered human acellular vessels
421 recellularize and evolve into living blood vessels after human implantation. *Sci.
422 Transl. Med.* **11**, 1–12.

423 8. L'Heureux, N. *et al.* (2006). Human tissue-engineered blood vessels for adult
424 arterial revascularization. *Nat. Med.* **12**, 361–365.

425 9. L'Heureux, N., McAllister, T. N. & de la Fuente, L. M. (2007). Tissue-
426 engineered blood vessel for adult arterial revascularization. *N. Engl. J. Med.*
427 **357**, 1451–1453.

428 10. L'Heureux, N. *et al.* (2007). Technology insight: the evolution of tissue-
429 engineered vascular grafts - from research to clinical practice. *Nat. Clin. Pract. Cardiovasc. Med.* **4**, 389–395.

430

431 11. Peck, M., Dusserre, N., McAllister, T. N. & L'Heureux, N. (2011). Tissue
432 engineering by self-assembly. *Mater. Today* **14**, 218–224.

433 12. Wu, W., Allen, R. A. & Wang, Y. (2012). Fast-degrading elastomer enables
434 rapid remodeling of a cell-free synthetic graft into a neoartery. *Nat. Med.* **18**,
435 1148–1153.

436 13. Cheng, S. *et al.* (2017). Self-Adjusting, polymeric multilayered roll that can
437 keep the shapes of the blood vessel scaffolds during biodegradation. *Adv. Mater.* **29**, 1700171.

438

439 14. Akentjew, T. L. *et al.* (2019). Rapid fabrication of reinforced and cell-laden
440 vascular grafts structurally inspired by human coronary arteries. *Nat. Commun.*
441 **10**, 1–15.

442 15. Hoenig, M. R., Campbell, G. R., Rolfe, B. E. & Campbell, J. H. (2005). Tissue-
443 engineered blood vessels: alternative to autologous grafts? *Arterioscler. Thromb. Vasc. Biol.* **25**, 1128–1134.

444

445 16. Yan, J., Lu, Y., Chen, G., Yang, M. & Gu, Z. (2018). Advances in liquid
446 metals for biomedical applications. *Chem. Soc. Rev.* **47**, 2518–2533.

447 17. Choi, S., Han, S. I., Kim, D., Hyeon, T. & Kim, D. H. (2019). High-
448 performance stretchable conductive nanocomposites: Materials, processes, and
449 device applications. *Chem. Soc. Rev.* **48**, 1566–1595.

450 18. Son, D. *et al.* (2015). Bioresorbable electronic stent integrated with therapeutic
451 nanoparticles for endovascular diseases. *ACS Nano* **9**, 5937–5946.

452 19. Du, Z. J., Bi, G. Q. & Cui, X. T. (2018). Electrically controlled neurochemical
453 release from dual-layer conducting polymer films for precise modulation of
454 neural network activity in rat barrel cortex. *Adv. Funct. Mater.* **28**, 1–12.

455 20. Gong, P. *et al.* (2013). A strategy for the construction of controlled, three-
456 dimensional, multilayered, tissue-like structures. *Adv. Funct. Mater.* **23**, 42–46.

457 21. Jin, Y. *et al.* (2013). Stress-induced self-assembly of complex three
458 dimensional structures by elastic membranes. *Small* **9**, 2410–2414.

459 22. Yuan, B. *et al.* (2012). A strategy for depositing different types of cells in three
460 dimensions to mimic tubular structures in tissues. *Adv. Mater.* **24**, 890–896.

461 23. Wang, N. *et al.* (2016). A strategy for rapid and facile fabrication of controlled,
462 layered blood vessel-like structures. *RSC Adv.* **6**, 55054–55063.

463 24. Tang, L., Mou, L., Zhang, W. & Jiang, X. (2019). Large-scale fabrication of
464 highly elastic conductors on a broad range of surfaces. *ACS Appl. Mater.*
465 *Interfaces* **11**, 7138–7147.

466 25. Tang, L. *et al.* (2018). Printable metal-polymer conductors for highly
467 stretchable bio-devices. *iScience* **4**, 302–311.

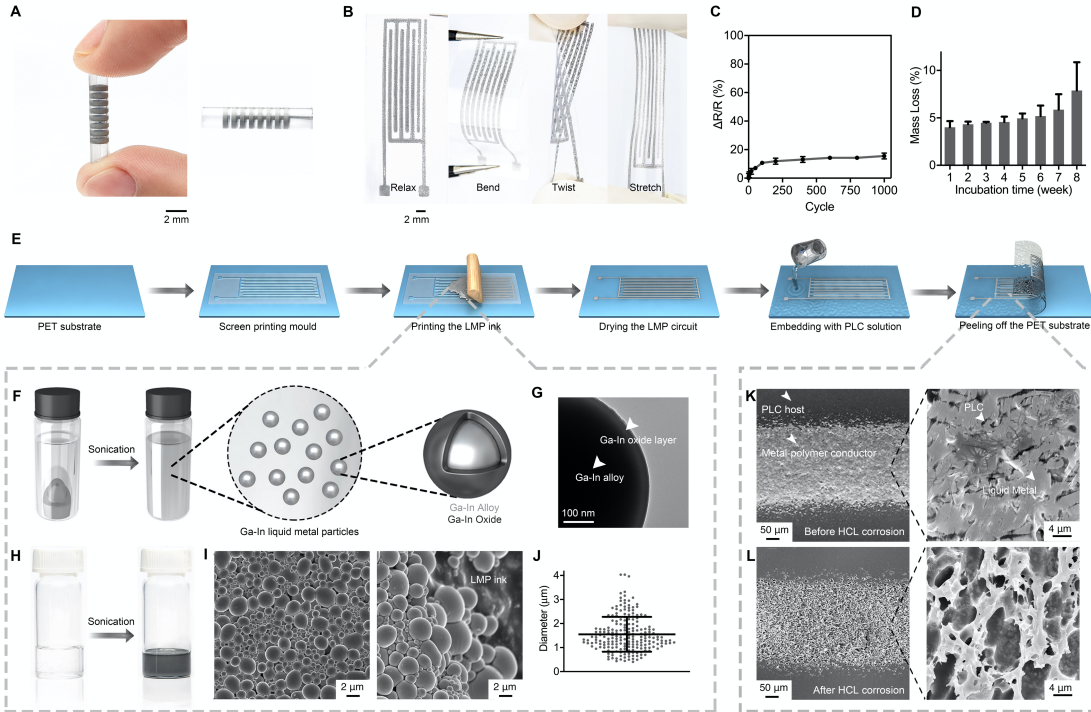
468 26. Bai, H., McCaig, C. D., Forrester, J. V. & Zhao, M. (2004). DC electric fields
469 induce distinct preangiogenic responses in microvascular and macrovascular
470 cells. *Arterioscler. Thromb. Vasc. Biol.* **24**, 1234–1239.

471 27. Lu, Y. *et al.* (2015). Transformable liquid-metal nanomedicine. *Nat. Commun.*

472 6, 10066.

473 28. Kenar, H. *et al.* (2019). Microfibrous scaffolds from poly(L-lactide-co- ϵ -
474 caprolactone) blended with xeno-free collagen/hyaluronic acid for
475 improvement of vascularization in tissue engineering applications. *Mater. Sci.*
476 *Eng. C* **97**, 31–44.

477



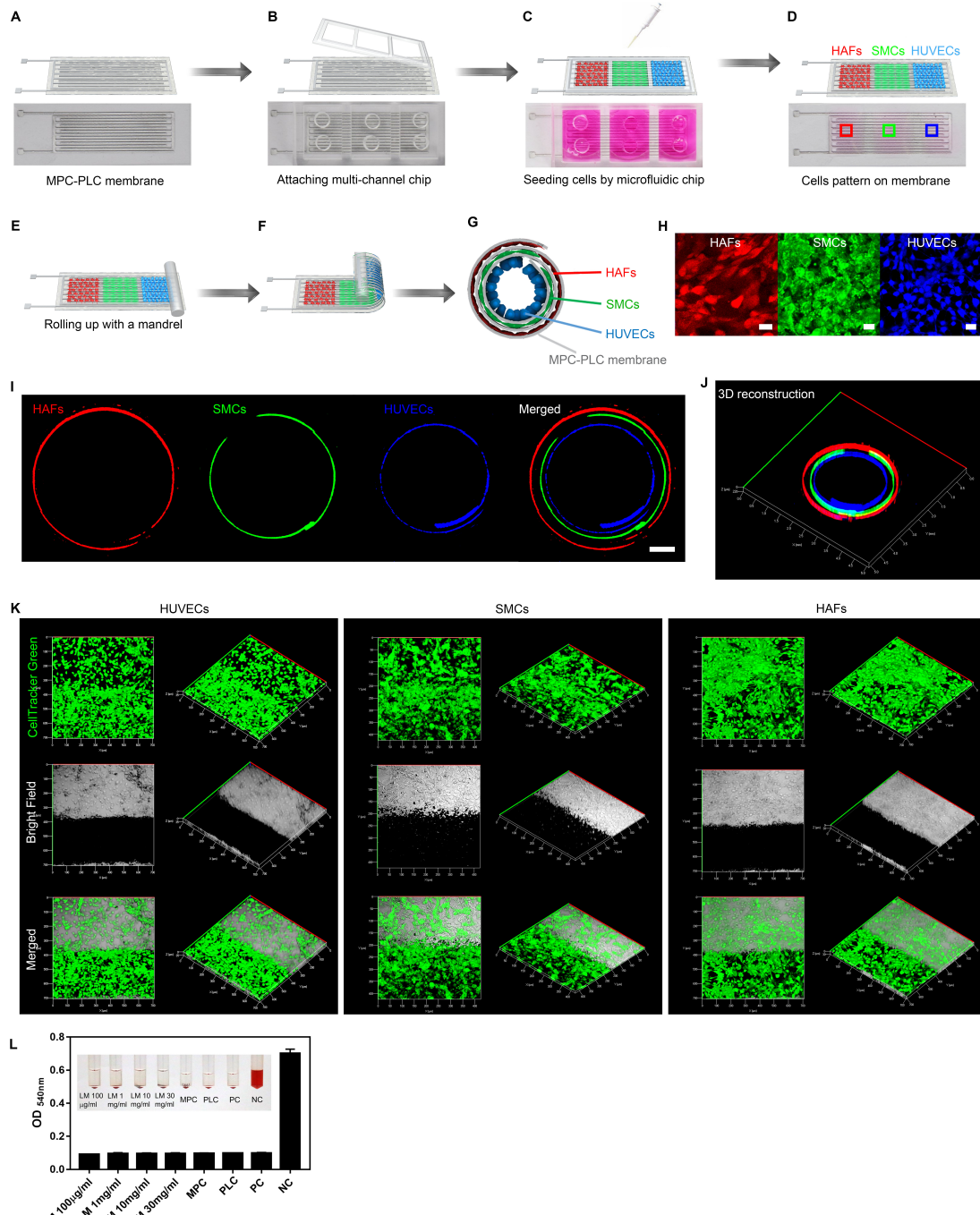
479 **Figure 1. Fabrication and characterization of the electronic blood vessel. (A)**

480 Snapshots of the electronic blood vessel. Scale bar, 4 mm. **(B)** Snapshots of the MPC-
481 PLC membrane. Scale bar, 1 mm. **(C)** $\Delta R/R$ changes with a bend of 180° for 1000
482 cycles (n=3). **(D)** Mass loss of the MPC-PLC membrane during 8-week incubation
483 (n=3 for each time point). **(E)** Schematic of fabrication of the MPC-PLC membrane.
484 **(F)** Schematic of fabrication of LMPs. **(G)** TEM image of the Ga-In particle. **(H)**
485 Photographs of mixture of the Ga-In alloy and the solvent, and the LMP ink after
486 sonication. **(I)** Representative SEM images of LMPs. Scale bar, 2 μ m. **(J)** Diameter
487 distribution of LMPs (n =200). **(K)** SEM images of the MPC-PLC circuit. White
488 arrowheads indicate the PLC host and MPC-PLC membrane. Scale bar, 50 μ m. Right
489 is the Zoom-in view. White arrowheads indicate the PLC and LMP ink. LMPs were
490 embedded by the PLC as the host polymer. Scale bar, 4 μ m. **(L)** SEM images of the
491 MPC-PLC circuit after corrosion by excessive hydrochloric acid. Scale bar, 50 μ m.

492 Right is the Zoom-in view. LMPs were fully removed by corrosion. The porous

493 structure only consisted of the PLC. Scale bar, 4 μm .

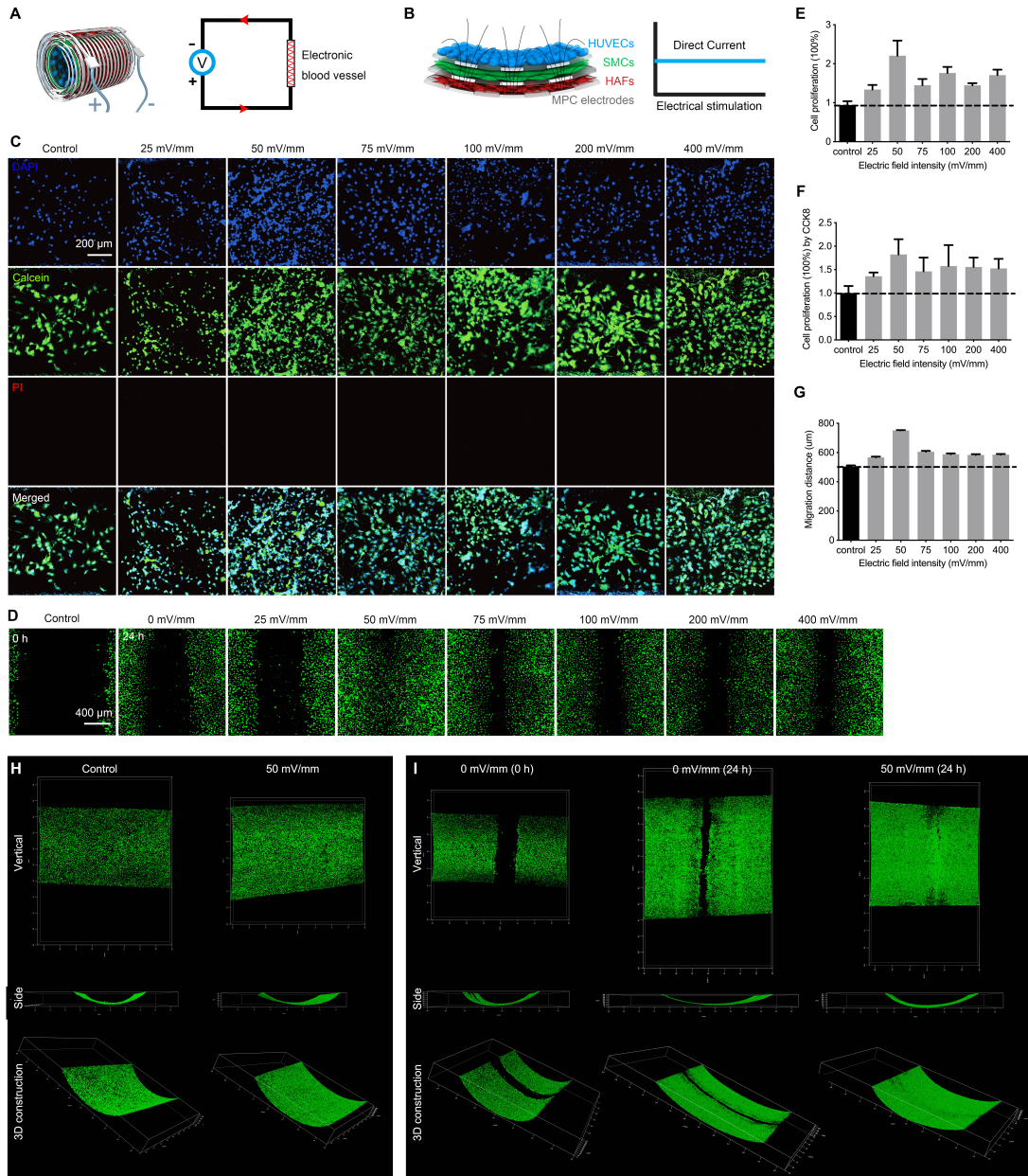
494



495

496 **Figure 2. *In vitro* biocompatibility of the electronic blood vessel. (A) The MPC-**
 497 **PLC membrane. (B) Attaching a multichannel microfluidic chip onto the MPC-PLC**
 498 **membrane. (C, D) Patterning three kinds of blood vessel cells (blue: HUVECs, green:**
 499 **SMCs, red: HAFs) onto the MPC-PLC membrane by using PDMS microfluidic chip.**
 500 **(E, F) Rolling the cell-laden membrane into a multi-layered tubular structure with a**

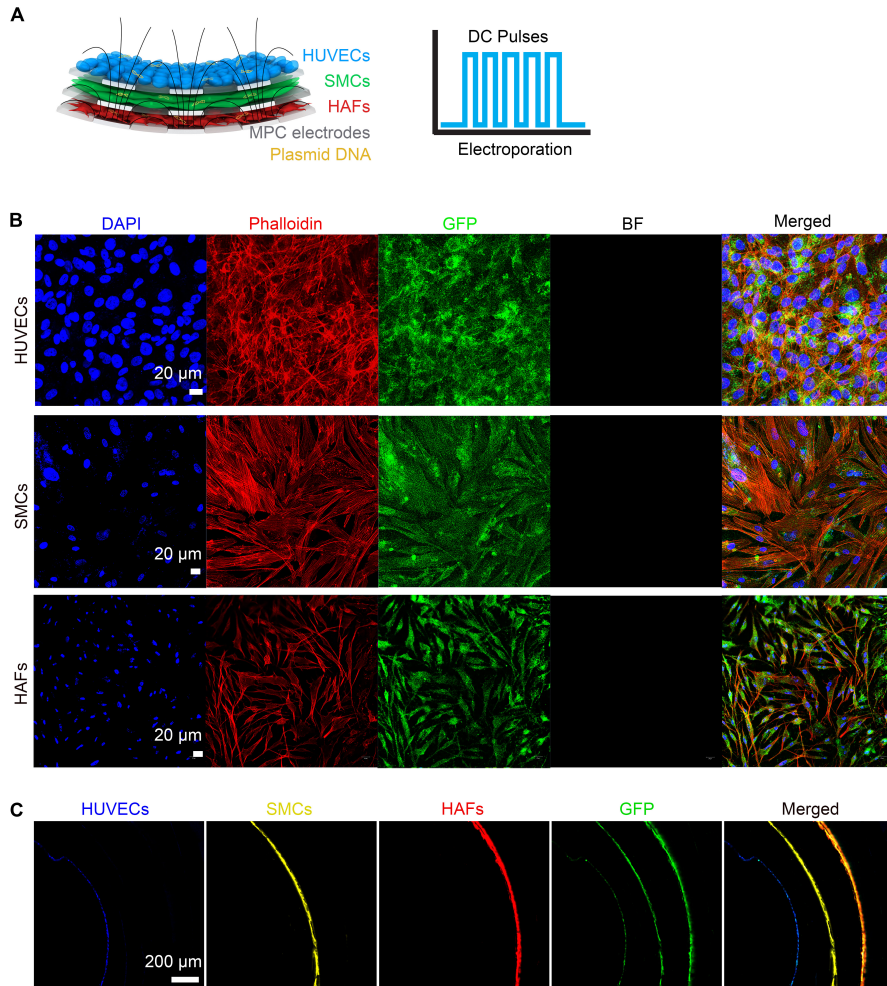
501 PTFE mandrel. **(G)** Cross-sectional view of natural blood vessel mimicking structure
502 with different cells distributed in different layers. **(H)** Fluorescent images of cells on
503 the MPC-PLC membrane, in corresponding to **(D)**. Stained by CellTracker violet,
504 green, deep red, respectively. Scale bar, 20 μm . **(I)** Three kinds of blood vessel cells
505 distributed in different layers of the electronic blood vessel. Scale bar, 500 μm . **(J)** 3D
506 construction of **(I)**. **(K)** Cell viability after 2-week incubation in the electronic blood
507 vessel. Stained by CellTracker green. The right column is the 3D reconstruction of the
508 left column in each image. The black area represents the MPC circuit and the
509 transparent area represents the PLC in bright-field images. **(L)** Hemolysis test. The
510 MPC circuit, PLC membrane, and liquid metal with different concentrations were
511 tested with the rabbit whole blood and exhibited good blood compatibility. Saline was
512 used as the positive control whereas water was used as the negative control.
513



514

515 **Figure 3. *In vitro* electrical stimulation. (A-B)** The electronic blood vessel was
516 connected to electrochemical station (A) to generate a direct current voltage (B). **(C)**
517 Confocal fluorescent images of HUVEC proliferation after 2 d incubation and
518 stimulation under different DC electric field: 25, 50, 75, 100, 200, 400 mV/mm,
519 respectively. Blue, DAPI; green, Calcein AM; red, PI. **(D)** Confocal fluorescent
520 images of HUVEC migration after 24 h incubation and stimulation under different

521 DC electric field: 25, 50, 75, 100, 200, 400 mV/mm, respectively. Stained by Calcein
522 AM. Scale bar, 400 μ m. A 10 μ l tip was used to scratch a line to create a model for
523 HUVECs migration. **(E)** The proliferation of HUVECs under different DC electrical
524 field, setting the control (without electric field) as 100%. (n=4). **(F)** The proliferation
525 of HUVECs under different DC electrical field tested with the CCK-8 kit. (n=4). **(G)**
526 The migration of HUVECs at different DC electrical field (n=4). **(H)** The
527 proliferation of HUVECs in a 3D model under DC electric field of 50 mV/mm.
528 Stained by Calcein AM. **(I)** The migration of HUVECs in a 3D model under DC
529 electric field of 50 mV/mm. Stained by Calcein AM. The HUVECs formed a
530 complete endothelial layer after 24 h electrical stimulation.
531



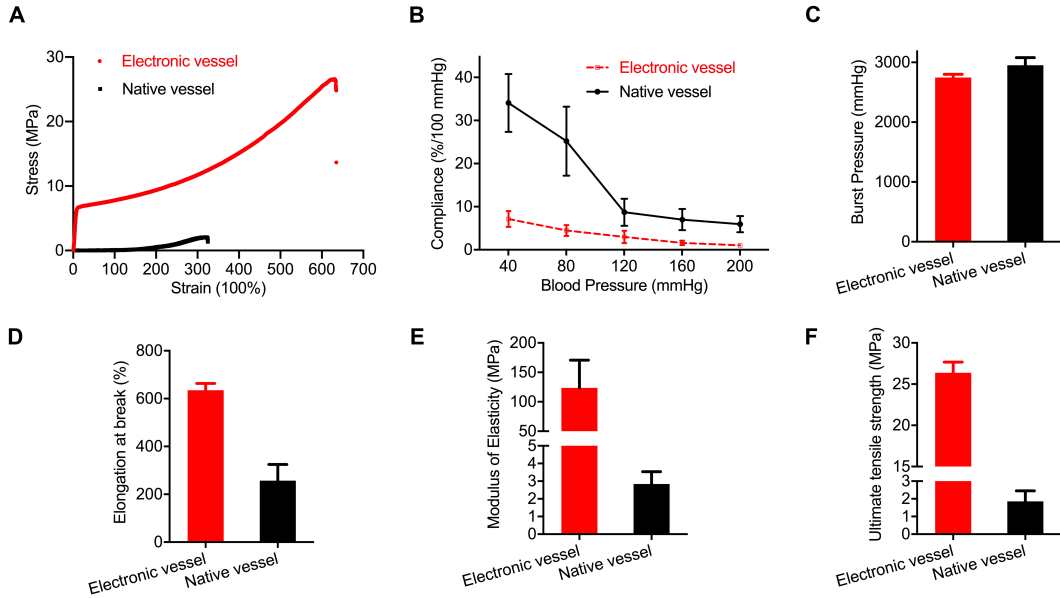
532

533 **Figure 4. *In vitro* electroporation. (A)** The electronic blood vessel was connected to
534 an electroporator to generate electrical pulses. **(B)** Confocal fluorescent images of
535 HUVECs, SMCs, and HAFs after electroporation under condition: voltage 60 V;
536 pulse duration 100 μ s; 5 pulses; pulse interval 1 s. Upper: HUVECs; middle: HAFs;
537 lower: SMCs. Blue represents cell nucleus, stained by DAPI; red represents cell
538 skeleton, stained by Rhodamine Phalloidin; green represents GFP; black represents
539 that cells were on the top of non-transparent MPC-PLC circuit, bright field. Scale bar,
540 20 μ m. **(C)** 3D distribution of HUVECs, SMCs, HAFs after electroporation under
541 condition: voltage 60 V; pulse duration 100 μ s; 5 pulses; pulse interval 1 s. Blue
542 represents HUVECs, stained by CellTracker Blue; yellow represents SMCs, stained

543 by CellTracker DiI; red represents HAFs, stained by CellTracker DiD; green

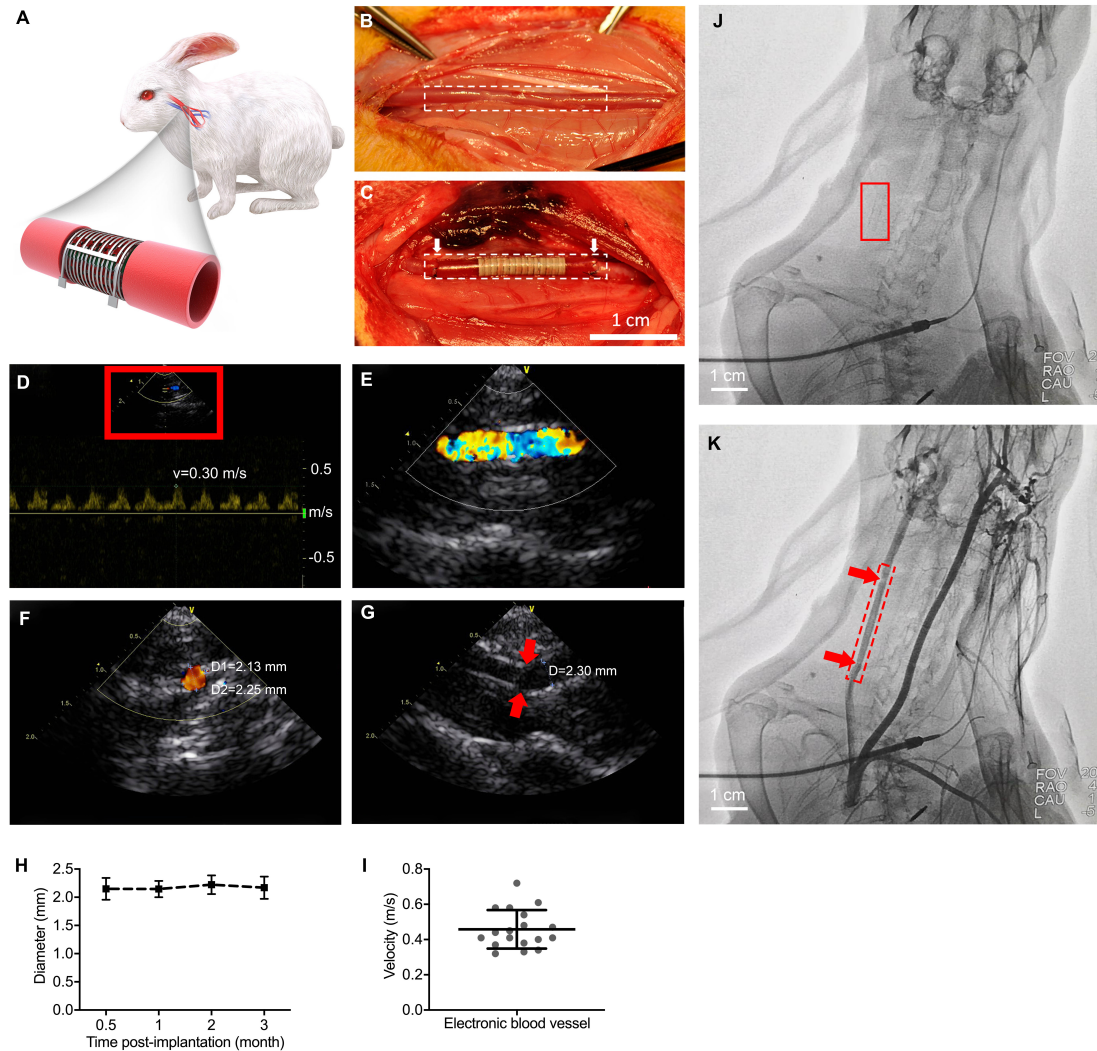
544 represents green fluorescent protein. Scale bar, 200 μ m.

545



546

547 **Figure 5. Mechanical properties of the electronic blood vessel. (A)** Stress-strain
548 curve. **(B)** Compliance test. **(C)** Burst pressure test. **(D)** Elongation at break. **(E)**
549 Modulus of elasticity. **(F)** Ultimate tensile strength. Electronic blood vessel (n= 5) and
550 native carotid artery (n= 3) were used in each test. All data are expressed as mean \pm
551 SD.
552

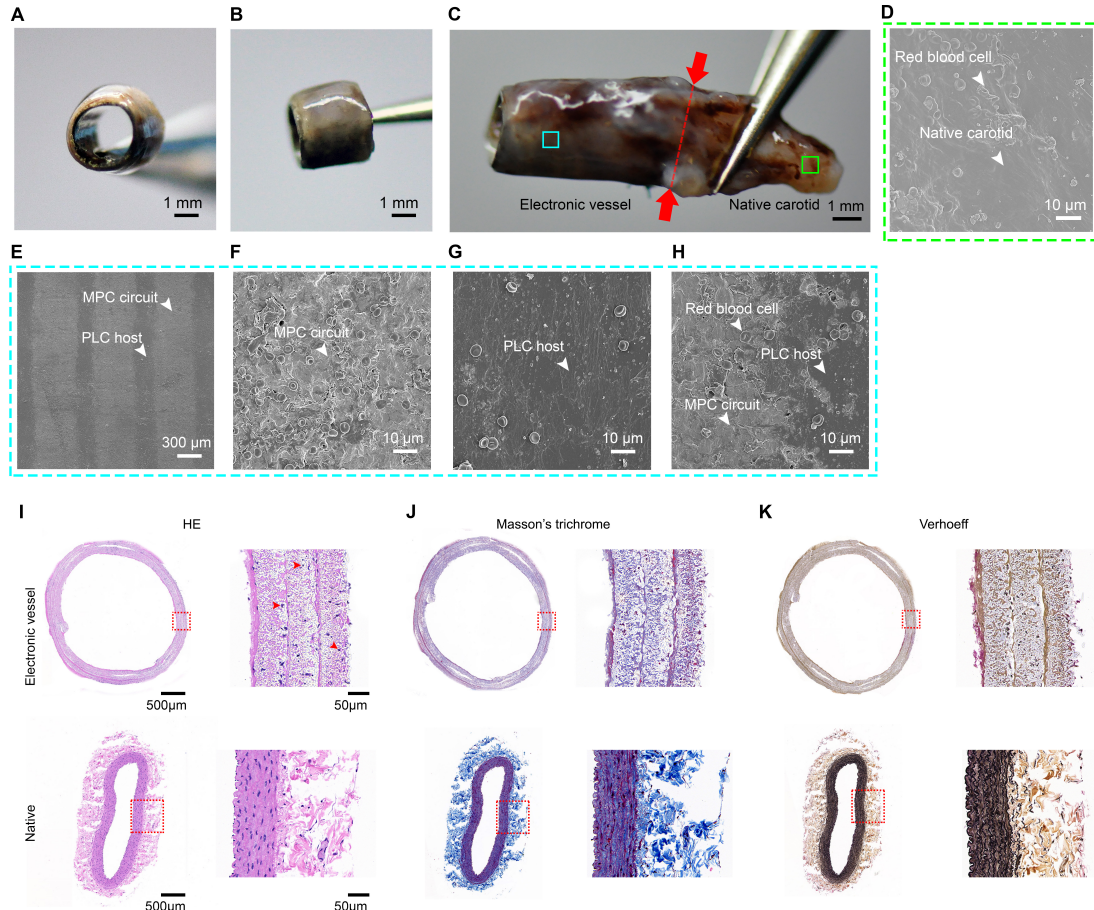


553

554 **Figure 6. *In situ* monitoring of the electronic blood vessel. (A)** Schematic of the
555 electronic blood vessel in the carotid artery of the rabbit. **(B, C)** An end-to-end
556 anastomosis procedure of electronic blood vessel implantation on carotid arteries of
557 rabbits (n=6). The dotted frames outline the margin of the native carotid artery and the
558 implanted electronic blood vessel. The white arrows indicate two ends of the
559 electronic blood vessel. Scale bar, 1 cm. **(D-G)** *In situ* monitoring of the electronic
560 blood vessel by doppler ultrasound imaging 3 months post-implantation.
561 Representative images from at least three different animals. **(D, E)** The real-time
562 blood flow at the operational site and the synchronized ultrasound pulses. The
563 asymmetric velocity curve represents the signal is from carotid artery rather than vein.
564 **(E)** Zoom-in view of red box in (D). **(F)** The cross view of the blood flow. **(G)** The
565 suture site (red arrows) connecting the native carotid artery and the electronic blood
566 vessel. **(H)** The diameter changes of the electronic blood vessel in different time post-

567 implantation. **(I)** The velocity of blood flow at the operational site (n=18, from
568 different rabbits at different time points). **(J, K)** *In situ* monitoring by arteriography 3
569 months post-implantation. **(J)** Image before injecting the contrast media. Red box
570 indicates the position of the implanted electronic blood vessel. **(K)** Image after
571 injecting the contrast media. Red box indicates the position of the implanted blood
572 vessel. Red arrows indicate the suture sites connecting the native carotid artery and
573 the electronic blood vessel.

574



575

576 **Figure 7. *Ex vivo* study of the electronic blood vessel after 3 months implantation**

577 **in the rabbit. (A-C)** The cross-sectional (A) and lateral view (B, C) of the explanted
578 electronic blood vessel after 3-month host remodeling. Red arrows and the dotted line
579 indicate the suture site connecting the native carotid artery (right) and the electronic
580 blood vessel (left). Scale bar, 1 mm. **(D)** Representative SEM image of the lumen of
581 the native carotid. Scale bar, 10 μ m. **(E-H)** Zoom-in view of the MPC
582 circuit (F), PLC host (G) and the interconnection area of MPC circuit and PLC host
583 (H). Scale bar, 10 μ m. **(I-K)** Hematoxylin/eosin (H&E) staining, Masson's trichrome
584 staining, and Verhoeff's staining of the implanted electronic blood vessel 3 months
585 staining, and Verhoeff's staining of the implanted electronic blood vessel 3 months

586 post-implantation, setting native carotid artery as control. The right panel is the zoom-
587 in view of the left panel. Scale bar (left), 500 μ m. Scale bar (right), 50 μ m.