Isochoric supercooled preservation and revival of human cardiac microtissues

Matthew J. Powell-Palm, Verena Charwat, Berenice Charrez, Brian Siemons, Kevin E. Healy & Boris Rubinsky

Low-temperature biopreservation and 3D tissue engineering present two differing routes towards eventual on-demand access to transplantable biologics, but recent advances in both fields present critical new opportunities for crossover between them. In this work, we demonstrate sub-zero centigrade preservation and revival of autonomously beating three-dimensional human induced pluripotent stem cell (hiPSC)-derived cardiac microtissues via isochoric supercooling, without the use of chemical cryoprotectants. We show that these tissues can cease autonomous beating during preservation and resume it after warming, that the supercooling process does not affect sarcomere structural integrity, and that the tissues maintain responsiveness to drug exposure following revival. Our work suggests both that functional three dimensional (3D) engineered tissues may provide an excellent high-content, low-risk testbed to study complex tissue biopreservation in a genetically human context, and that isochoric supercooling may provide a robust method for preserving and reviving engineered tissues themselves.
Modem organ and tissue transplantation efforts in the U.S. and abroad are hamstrung by the unavailability of on-demand donor biologics1–3. While this may be interpreted in part as a matter of policy, the central scientific driver of this organ and tissue deficit is the extremely short ex vivo shelf life of complex biologics. In the U.S. alone, though more than 100,000 people are currently on transplant waiting lists, more than 70% of all thoracic donor organs are discarded each year, simply due to an inability to preserve them for sufficient time periods4. The preservable ex vivo lifetime of donor hearts for instance, which are typically held on ice in "static cold storage", is only 4–6 h1.

While historical research on the preservation of organs and complex tissues focused largely on the deep cryogenic temperatures that would theoretically enable indefinite storage, the last decade has seen the advent of "high-subzero" preservation efforts, targeting the −20°–0°C range5. These efforts aim simply for multi-day preservation, in recognition of the fact that even 24- to 48-h preservation of many complex donor biologics could prove transformative for transplant medicine1–4. Chief amongst high-subzero efforts stands supercooled storage, which aims, via a number of different protocols, to hold biologics in an ice-free, metastable liquid preservation solution at subzero temperatures6–9.

In this work, we employ a recently developed technique, isochoric supercooling6, to achieve high-stability and predictable supercooling of the University of Wisconsin (UW) organ preservation solution at −3°C, in which we successfully preserve a functional 3D hiPSC-derived cardiac microphysiological system (MPS)10 for multiple days without the addition of non-physiological cryoprotectants such as dimethyl sulfoxide (DMSO) or glycerol.

Results

Cardiac MPS supercooled preservation protocol. The cardiac MPS combines human cells with microfluidics (Fig. 1a, b) to promote 3D self-assembly into a microtissue (Fig. 1c) that faithfully recapitulates complex human heart muscle structure and function10–14. The cardiac microtissues used in this work express a genetically encoded fluorescent calcium channel reporter10, a convenient means to optically and non-invasively track beating activity via calcium flux (Fig. 1d). The calcium transients can be analyzed to obtain important information on beating activity such as beat rate, beat duration (in this study we use calcium waveform as a proxy for action potential duration (APD)11), and aberrations in beating. The MPS also allows for external electrical tissue stimulation via pacing electrodes (Fig. 1d).

In designing the proof-of-concept preservation effort, we established the following criteria as indicators of successful preservation: retention of sarcomere structural integrity; retention of physiological function including autonomous beating activity and responsiveness to external electrical pacing stimuli; and retention of responsiveness to a pharmacological inotrope (here the β-adrenergic agonist, isoproterenol). Not only were these metrics rapidly monitored in the MPS, they can also be easily adapted to assess the preservation success of native heart tissue or full organs in a clinical setting (e.g., ECG, QT interval).

Inspired by previous supercooled preservation efforts involving the liver7,8, we designed a temperature–time trajectory by which to shuttle the cardiac MPS into and out of supercooled preservation. This trajectory is shown in Fig. 1f, and includes the following essential steps: replacing all metabolizable media in the MPS with non-metabolizable saline solution (PBS, no calcium); submerging the MPS in a bath of PBS initially at 37°C, which is then cooled to 1 ± 1°C at a rate of −2°C/min; replacing the PBS in the MPS with UW organ preservation solution (pre-chilled to 1 ± 1°C); placing the MPS in a rigid isochoric container filled with pre-chilled UW organ preservation solution; sealing the container tightly without trapping any bulk gas phase6 (Fig. 1c); and, finally, submerging the isochoric chamber in a constant-temperature circulating cooling bath at −3°C for the target preservation duration (24, 48, or 72 h). Per Fig. 1f, these steps are performed in reverse for rewarming after preservation. After the MPS are returned to 37°C and their supply of metabolizable media is replenished, they are imaged once without pacing, then moved to a cell culture incubator and allowed a minimum of 24 h of recovery before further imaging, pacing, and application of isoproterenol. A more detailed accounting of the protocol is available in the Methods section.

Cardiac MPS supercooled preservation outcomes. The results of this preservation process are presented in Fig. 2. An average of 50–65% of tissues resumed normal spontaneous beating activity after isochoric supercooling (65% after 24-h preservation; 50% after 48-h preservation; 55% after 72-h preservation), and a similar percentage (50–70%) responded normally to external electrical stimulation (Fig. 2a), with no statistically significant differences observed between preservation durations. An additional 5–15% of MPS yielded the partial recovery of the tissue (gray bars in Fig. 2a), and the remaining 20–40% did not resume coherent beating. The explicit mechanistic drivers at play are not yet known and must be probed in future studies, but it should be noted that failure to resume coherent beating does not necessarily imply cell death—tissue beating is a combination of complex electrophysiological and mechanical processes reliant on myriad intra- and intercellular behavior, the corrosion of any of which could contribute to failure.

Immunofluorescence staining confirmed qualitatively that the bulk structural integrity of the tissues was not altered by the cooling and warming processes, as shown by comparable post-preservation size and distribution of nuclei, as well as general alignment of cardiomyocytes and sarcomeric integrity (Fig. 2b and Supplementary Fig. S1). Further evaluation of physiological properties revealed that neither beat rate nor beat shape (as measured by a triangulation metric (APD80–APD30)/APD8015) was significantly altered by isochoric supercooling (Fig. 2c, e). Beat duration (APD80) showed a slight increasing trend with increasing supercooling both in spontaneous and externally paced conditions, though any potential biological relevance of this trend (Fig. 2d) is not apparent and should be explored in more detail in future studies.

In order to probe the conservation of drug response characteristics, we subjected the post-preservation MPS to 1μM of the well-known β-adrenoceptor agonist isoproterenol. The observed increase in spontaneous beat rate further confirmed the recovery of essential cardiac tissue function after isochoric supercooling, though the average beat rate increased by approximately 40% (Fig. 2f), which is less than previously reported for MPS created from the same hiPSC line12. It is unclear however whether this difference is a product of the supercooling process or a simple result of batch-to-batch variation within the microtissues, as the present tissues were not exposed to isoproterenol before supercooling (in order to avoid potentially confounding effects of pre-preservation drug exposure). The potential between supercooled preservation and altered drug response should be investigated further in future work.

With the exception of the slightly increasing trend in APD80, all recovery rates, electrophysiological parameters, and drug
response characteristics did not vary significantly with supercooling duration (24, 48, or 72 h), suggesting that the mechanisms of tissue damage are not sharply dependent on metabolism and that significantly longer preservation periods may be achievable with similarly high recovery. We hypothesize that the majority of tissue damage occurs during the cooling/warming processes, which are independent of supercooling duration and were not fully optimized in this study. Excessive cooling rates may produce damaging stresses in the tissue due to a mismatch between the thermal contractility of the tissue and the surrounding microchamber. Damage may also be driven by the failure to sufficiently remove metabolizable or ionic constituents prior to cooling, which can generate toxicity during rewarming due to ionic imbalances or accumulation of metabolic byproducts. It should also be noted that the observed failure of some MPS to resume beating may be a product of the inherent variability in the MPS themselves, which is evident in the pre-supercooling violin plots shown in Fig. 2c–e, and that this variability may mask smaller effects within the investigated parameter space.

Importantly, we observe no significant morphological or electrophysiological changes in recovered tissues that would suggest corruption of core cardiac functionality, as embodied most essentially by the resumption of spontaneous autonomous beating with an unchanged beat shape and expected responsiveness to isoproterenol treatment.

Discussion

Our data demonstrate that functional 3D microtissues may be effectively preserved at subzero temperatures, which has several important implications.

First, these microtissue constructs may offer essential translational insights into expected physiological effects of low-temperature preservation at the whole-organ scale. Indeed, in the drug response and pharmacokinetic analysis domains, various human-derived MPS have been demonstrated to be superior predictors of clinical response compared to animal models, offering closer physiologic and pharmacokinetic properties to native tissue whilst enabling higher-throughput testing. Furthermore, recent work has demonstrated that the cardiac MPS can successfully predict clinical trial arrhythmia outcomes, indicating potential prognostic value in predicting human-scale cardiac behavior. We suggest that these same advantages can apply to low-temperature preservation, and that MPS may provide a better testbed for identifying the missing links required for timely translation of benchtop preservation protocols to the clinic, where advances in organ preservation are urgently needed. Translatability to full-organ systems will of course require computational analysis of thermal/mass transport and metabolic scaling issues, as well as accounting for the fundamental differences in structural and electrophysiological complexity between the MPS and the native system. However, the complex functional...
outputs that can be probed with the MPS may move benchtop preservation research a significant step closer to clinical relevance.

Second, the high-recovery preservation results obtained using the protocol described herein, which does not include the introduction of non-physiological cryoprotectants, suggest that MPS platforms may also be used to study the explicit effects of various cryoprotectants on preservation efficacy. MPS platforms could be used to simultaneously screen cryoprotectants for toxicity (in a complex genetically human context), cryoprotective action, and diffusion/perfusion performance. The use of non-physiological cryoprotectants has been omnipresent in cellular preservation protocols since the 1950s, but has failed to penetrate clinical full-organ preservation, largely due to concerns surrounding toxicity or perfusion in native human tissue, and poor scaling of cell preservation results. Our data establish a baseline from which MPS may be used for higher-throughput, higher-reliability cryoprotectant screening.

This work also, to the best of our knowledge, presents the first demonstration of biological preservation by isochoric supercooling. This technique provides a thermodynamically simple and procedurally streamlined method of ice-free preservation at sub-0 °C temperatures, and given its non-reliance on specific cryoprotective agents, we suggest that it may be applied for the preservation of arbitrary biologics in the high-subzero temperature range.

Fig. 2 Recovery rate and comparison of key cardiac functionality parameters pre- and post-isochoric supercooling. Statistical difference between groups (p < 0.05) is marked by horizontal lines with asterisks. All raw data points are shown within violin plots. a Total recovery rate of preserved tissues as a function of preservation time. Gray bars indicate tissues that presented some activity post-isochoric supercooling, but were not coherent over the entire tissue (i.e., only part of the microtissue recovered). Scatter markers represent the weighted means of individual experiments, and marker diameter reflects the number of tissues (total sample sizes, i.e., number of independent cardiac tissues, for each condition: 24 h: n = 39; 48 h: n = 36; 72 h: n = 17). Error bars mark the standard error of the mean (SEM) across experimental groups, weighted by the total number of tissues evaluated in each group. These error bars reflect the SEM of the total surviving tissue population, i.e., the combined population of fully recovered and partially recovered tissues.

b Confocal microscopy image of cardiac microtissue before and after 24 h supercooled preservation (α-actinin in red and nuclei in blue). c Triangulation ((APD80-APD30)/APD80) as a metric of beat shape following 24, 48, or 72 h isochoric supercooling. All raw data points are scatter plotted within the violin plots. d Percent change in beat rate-corrected APD80 of recovered tissues compared to pre-isochoric supercooling. e Spontaneous beat rate of recovered tissues. f Increase in normalized beat rate upon introduction of 1 μM isoproterenol for 30 min to tissues that had previously been subjected to 24 h (left) or 72 h (right) supercooled preservation (sample sizes: 24 h: n = 6; 72 h: n = 5).
Building upon the baseline data, future work may probe a wide variety of additional complex functional parameters available from the cardiac MPS. Perhaps most pressing, the contractile force and relaxation should be analyzed post-supercooling, which can provide a more definitive measure of health and function in the revived tissues. Additional analyses may include metabolic activity, biomarker secretion, or PCR studies.

In summary, this study demonstrates successful isochoric supercooled preservation and recovery of microfluidic human cardiac tissue cultures based on structural and functional characteristics. Further molecular biological studies will be needed to assess any changes in gene and biomarker protein expression, and optimization of thermal cycling protocols should be explored to maximize recovery rate. Moving forward, as MPS platforms continue to advance in both fidelity and accessibility, we suggest that MPS preservation studies may provide a useful preliminary screening tool for any full-organ or complex tissue preservation efforts, providing insight into potential structural and functional effects of the preservation process on high-content, complex and human-derived tissues.

Methods

Preservation—cooling. The cooling protocol follows the trajectory plotted in Fig. 1. Each chip, on a 37 °C hotplate (Tokai Hit, Gendoji-cho, Japan), was first washed with 70% ethanol to reduce contamination risks. The MPS were then transferred to an approximately 65 mL thick-walled glass container (Corning, 354277) in mTeSR-1 media (Stem Cell Technologies, 85851) for several seconds. Each chip was then directly placed into a circulating ice and water bath at 1 ± 1 °C, resulting in a cooling rate of 2.1 ± 0.15 °C/min. The temperature in both the well and a reference MPS were continuously monitored via a thermocouple. The preservation temperature remained constant at 37 °C. This well was then placed directly into a circulating ice and water bath at 1 ± 1 °C (via ice bath) before it was opened. Premature opening of the isochoric preservation and warming in order to minimize the risk of calcium shock or other ionic activity, biomarker secretion, or PCR studies.

CM differentiation from iPSC. Cell culture was performed similarly to already published protocols12. We used the human iPSC line Wild Type C (WTC; #GM25256, Coriell Institute) edited to express a fluorescent reporter for intracellular calcium (GCaMP6F)28. After thawing the iPSCs were cultured on Matrigel (Corning, 354277) in mTeSR-1 media (Stem Cell Technologies, 85851) for several passages until every 3 days media was changed in parallel. Media channels were centrifuged off the master mold and cut to size. Holes of 0.75 mm diameter were punched out of each master mold and placed in parallel on a microfluidic chip. Media exchange in the cell chamber was achieved via an array of small connection channels (2 × 2 μm cross-section; 40 μm length) that protected the tissue from fluid mechanical forces31. Small pillars (20 μm diameter) at either end of the cell chamber served as structural anchor points to keep the muscle fiber from collapsing. Each microfluidic device featured either a single tissue chamber or 4 identical tissue chambers in parallel.

CM differentiation from iPSC. Cell culture was performed similarly to already published protocols12. We used the human iPSC line Wild Type C (WTC; #GM25256, Coriell Institute) edited to express a fluorescent reporter for intracellular calcium (GCaMP6F)28. After thawing the iPSCs were cultured on Matrigel (Corning, 354277) in mTeSR-1 media (Stem Cell Technologies, 85851) for several passages until every 3 days media was changed in parallel. Media channels were centrifuged off the master mold and cut to size. Holes of 0.75 mm diameter were punched out of each master mold and placed in parallel on a microfluidic chip. Media exchange in the cell chamber was achieved via an array of small connection channels (2 × 2 μm cross-section; 40 μm length) that protected the tissue from fluid mechanical forces31. Small pillars (20 μm diameter) at either end of the cell chamber served as structural anchor points to keep the muscle fiber from collapsing. Each microfluidic device featured either a single tissue chamber or 4 identical tissue chambers in parallel.

CM differentiation from iPSC. Cell culture was performed similarly to already published protocols12. We used the human iPSC line Wild Type C (WTC; #GM25256, Coriell Institute) edited to express a fluorescent reporter for intracellular calcium (GCaMP6F)28. After thawing the iPSCs were cultured on Matrigel (Corning, 354277) in mTeSR-1 media (Stem Cell Technologies, 85851) for several passages until every 3 days media was changed in parallel. Media channels were centrifuged off the master mold and cut to size. Holes of 0.75 mm diameter were punched out of each master mold and placed in parallel on a microfluidic chip. Media exchange in the cell chamber was achieved via an array of small connection channels (2 × 2 μm cross-section; 40 μm length) that protected the tissue from fluid mechanical forces31. Small pillars (20 μm diameter) at either end of the cell chamber served as structural anchor points to keep the muscle fiber from collapsing. Each microfluidic device featured either a single tissue chamber or 4 identical tissue chambers in parallel.

CM differentiation from iPSC. Cell culture was performed similarly to already published protocols12. We used the human iPSC line Wild Type C (WTC; #GM25256, Coriell Institute) edited to express a fluorescent reporter for intracellular calcium (GCaMP6F)28. After thawing the iPSCs were cultured on Matrigel (Corning, 354277) in mTeSR-1 media (Stem Cell Technologies, 85851) for several passages until every 3 days media was changed in parallel. Media channels were centrifuged off the master mold and cut to size. Holes of 0.75 mm diameter were punched out of each master mold and placed in parallel on a microfluidic chip. Media exchange in the cell chamber was achieved via an array of small connection channels (2 × 2 μm cross-section; 40 μm length) that protected the tissue from fluid mechanical forces31. Small pillars (20 μm diameter) at either end of the cell chamber served as structural anchor points to keep the muscle fiber from collapsing. Each microfluidic device featured either a single tissue chamber or 4 identical tissue chambers in parallel.
The cell loading port was then sealed and E820 + RI media was perfused through the media channels after 1 h incubation time. The next day media was exchanged to RPMI ± 2% B27 + C and changed three times per week thereafter. MPS were used within 30 days after loading.

**Data acquisition.** Videos were obtained using a digital CMOS camera (HAMAMATSU, C11440/ORCA-Flash 4.0) mounted to Nikon TE-300 inverted microscope with an LED light engine (Lumencor SpectraX). A heated platform (Tokai Hit, TPF SQQ) was used to image the MPS at 37 °C. For electrical stimulation, a pulse generator (ION OPTIX Myopacer Field Simulator) was used. 20 V biphasic rectangular pulses (20 ms) were applied via 1.5” long blunt stainless-steel needles (Vita Needle M937) inserted into the media inlet and outlet pipette tips. Calcium traces were recorded at 100 fps with 4 × 8 binning using the cyan LED for excitation. A 1 mM isoproterenol stock was prepared freshly from isoproterenol hydrochloride (J0260; TCI Chemicals) powder right before the drug response experiments, sterile filtered and diluted to 1 μM. 100 μL of drug-containing media were applied to each chip via the inlet pipette tip 30 min before recording.

**Data analysis.** Fluorescence videos were converted into time-intensity profiles (for example Fig. 1d) using an in-house library of Python scripts. Beat parameters such as beat rate and peak duration at 30% and 80% intensity (APD30 and APD80, respectively) were automatically detected. Further metrics were calculated: The beat rate-corrected peak duration (APD90) was computed using the Frankelid method, which corrects for variations in beat rate by scaling the values to 1 Hz beat rate; EPr = APDp/BR 1/2 with BR being the beat rate in Hz and cAPD90 being the peak duration at 80% peak height corrected to a beat rate of 1 Hz. Triangulation, as introduced by Hondegem et al.15, was computed as a metric of beat shape: (APDp – APD90)/APDac. Lower triangulation values indicate a more rectangular beat shape while higher values indicate stronger triangulation, which is associated with increased pro-arrhythmic potential.

For analysis of the recovery rate, we counted the percentage of tissues that resumed beating activity after preservation as judged by contractile tissue motion or peaks in the calcium trace. We distinguished between samples that showed coherent beating of the entire tissue as one unit from those that showed activity only in a portion of the tissue. We defined recovery rate as the number of tissues that resumed beating activity spontaneously or in response to electrical stimulation at any time point after preservation, divided by the total number of preserved tissues.

**Statistics and reproducibility.** All graphs were plotted in MATLAB (version R2019b). Violin plots were generated using a third-party MATLAB script developed by Bastian Bechtold (available here: https://github.com/bastibe/Violinplot-Matlab). Figures were assembled using Adobe Illustrator (version 25.0.1). Statistical analysis to compare preservation effect was done by one-way ANOVA, with statistical significance levels of p < 0.05 (*). In the presented data, horizontal bars topped by an asterisk indicate a significance level of p < 0.05 (*). The presented data are all available upon reasonable request. All data points are plotted in Figs. 1 and 2 are available in the Supplementary Data 1 file.

**References**

1. Giwa, S. et al. The promise of organ and tissue preservation to transform medicine. Nat. Biotechnol. 35, 520–542 (2017).
2. Brutsaert, B. G. & Uygur, H. Human heart preservation: The dawn of a new ice age? Curr. Opin. Organ Transplant. https://doi.org/10.1097/MOT.0000000000000403 (2017).
3. Fuller, B. J., Pettenko. A. & Guibert, E. Human organs come out of the deep cold. Nat. Biotechnol. https://doi.org/10.1038/nbt.3948-019-0264-2 (2019).
4. Ward, A., Klassen, D. K., Franz, K. M., Giwa, S. & Lewis, J. K. Social, economic, and policy implications of organ preservation advances. Curr. Opin. Organ Transplant. https://doi.org/10.1097/MOT.0000000000000532 (2018).
5. Taylor, M. J., Weegman, B. P., Baius, S. C. & Giwa, S. E. New approaches to cryopreservation of cells, tissues, and organs. Transfus. Med. Hemother. https://doi.org/10.1159/000499453 (2019).
6. Hordnell, M. J., Koh-Bell, A. & Rubinsky, B. Ischordic conditions enhance stability of metastable supercooled water. Appl. Phys. Lett. https://doi.org/10.1063/1.5143354 (2020).
7. Berendsen, T. A. et al. Supercooling enables long-term transplantation survival following 4 days of liver preservation. Nat. Med. https://doi.org/10.1038/nm.3588 (2014).
8. de Vries, R. J. et al. Supercooling extends preservation time of human livers. Nat. Biotechnol. https://doi.org/10.1038/s41587-019-0223-y (2019).
9. Huang, H., Yarmush, M. L. & Usta, O. B. Long-term deep-supercooling of large-volume water and red cell suspensions via surface sealing with immiscible liquids. Nat. Commun. https://doi.org/10.1038/s41467-018-05636-0 (2018).
10. Mathur, A. et al. Human iPSC-based cardiac microphysiological system for drug screening applications. Sci. Rep. 5, 8883 (2015).
11. Charrez, B. et al. In vitro safety “clinical trial” of the cardiac liability of drug polytherapy. Clin. Transl. Sci. https://doi.org/10.1111/cts.13038 (2021).
12. Huebsch, N. et al. Metabolically-driven maturation of hiPSC-Cell derived heart-on-a-chip. Preprint at bioRxiv https://doi.org/10.1101/485169 (2018).
13. Jæger, K. H. et al. Improved computational identification of drug response using optical measurement of human stem cell derived cardiomyocytes in microphysiological systems. Front. Pharmacol. 10, 1648 (2020).
14. Tveito, A. et al. Inversion and computational maturation of drug response using human stem cell derived cardiomyocytes in microphysiological systems. Sci. Rep. 8, 17626 (2018).
15. Hondegem, L. M., Carlson, L. & Duker, G. Instability and triangulation of the action potential predict serious proarrhythmia, but action potential duration prolongation is antiaarrhythmic. Circulation 103, 2004–2013 (2001).
16. Rubinsky, B. Principles of low temperature cell preservation. Heart Fail. Rev. 8, 277–284 (2003).
17. Ingber, D. E. Is it time for reviewer 3 to request human organ chip experiments instead of animal validation studies? Adv. Sci. 7, 2002030 (2020).
18. Ma, C., Peng, Y., Li, H. & Chen, W. Organ-on-a-chip: a new paradigm for drug development. Trends Pharmacol. Sci. 42, 119–133 (2021).
19. Mittal, R. et al. Organ-on-chip models: Implications in drug discovery and clinical applications. J. Cell. Physiol. 234, 8352–8380 (2019).
20. Faby, G. M. The relevance of cryoprotectant “toxicity” to cryobiology. Cryobiology https://doi.org/10.1006/cbio.1996.0011 (1996).
21. Zhang, B., Koroli, A., Lai, B. F. L. & Radisic, M. Advances in organ-on-a-chip engineering. Nat. Rev. Mater. 3, 257–278 (2018).
22. Wu, Q. et al. Organ-on-a-chip: Recent breakthroughs and future prospects. BioMedical Eng. Online 19, 9 (2020).
23. Powell-Palm, M. J., Koh-Bell, A. & Rubinsky, B. Ischordic conditions enhance stability of metastable supercooled water. Appl. Phys. Lett. https://doi.org/10.1063/1.5101118 (2021).
24. Huang, H., Yarmush, M. L. & Usta, O. B. Long-term deep-supercooling of large-volume water and red cell suspensions via surface sealing with immiscible liquids. Nat. Commun. https://doi.org/10.1038/s41467-018-05636-0 (2018).
25. Consiglio, A., Utkapi, G., Rubinsky, B. & Powell-Palm, M. J. Suppression of cavitation-induced nucleation in systems under isochoric confinement. Phys. Rev. Res. 2, 23350 (2020).
26. Charrez, B. et al. In vitro safety “clinical trial” of the cardiac liability of drug polytherapy. Clin Transl Sci. 14, 1155–1165 (2021).
27. Gohil, S. V., Suhail, S., Rubinsky, B. & Nair, L. S. Polymers and composites for orthopedic applications. Mater. Devices Bone Disord. https://doi.org/10.1007/s12079-012-00008-2 (2017).
28. Huebsch, N. et al. Automated video-based analysis of contractility and calcium flux in human-induced pluripotent stem cell-derived cardiomyocytes cultured over different spatial scales. Tissue Eng. C Methods 21, 467–479 (2015).
29. Vandenberk, B. et al. Which QT correction formulae to use for QT monitoring? J. Am. Heart Assoc. 5, e003264 (2016).
30. Powell-Palm, M. J., Rubinsky, B. & Sun, W. Freezing water at constant volume and under confinement. Commun. Phys. https://doi.org/10.1038/s42005-020-0303-9 (2020).
31. Consiglio, A., Ukpai, G., Rubinsky, B. & Powell-Palm, M. J. Suppression of cavitation-induced nucleation in systems under isochoric confinement. Phys. Rev. Res. https://doi.org/10.1103/physrevresearch.2.023350 (2020).

Acknowledgements
Funding is gratefully acknowledged from the NSF Engineering Research Center for Advanced Technologies for Preservation of Biological Systems (ATP-Bio) NSF EEC #1941543. We would also like to thank Dr. Henrik Finsberg and Dr. Sam Wall (Simula Research Laboratory, Oslo, Norway) for the Python scripts used to extract beat information from calcium traces, Dr. Bruce Conklin (Gladstone Institutes, San Francisco, USA) for providing the hiPSCs and technical advice on the cell line, and Dr. Chenang Lyu for technical assistance with the isochoric chamber assembly process.

Author contributions
M.J.P.-P., B.R. and K.E.H. conceived the study. M.J.P.-P., V.C., B.C. and B.A.S. designed the preservation protocol, conducted all experiments, collected and analyzed all data, and wrote the manuscript. B.R. and K.E.H. reviewed the data and edited the manuscript.

Competing interests
K.E.H. and B.A.S. have a financial relationship with Organos Inc. and both they and the company may benefit from the commercialization of the results of this research. M.J.P.-P. and B.R. have a financial relationship with BioChoric Inc. and both they and the company may benefit from the commercialization of the results of this research. The remaining authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s42003-021-02650-9.

Correspondence and requests for materials should be addressed to Matthew J. Powell-Palm, Kevin E. Healy or Boris Rubinsky.

Peer review information Communications Biology thanks Bramasta Nugraha, Katja Schenke-Layland and the other, anonymous, reviewers for their contribution to the peer review of this work. Primary Handling Editor: Eve Rogers.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2021