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| 1 | Base Editing Correction of Hypertrophic Cardiomyopathy in Human |
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| 2 | Cardiomyocytes and Humanized Mice |
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14 Abstract

The most common form of genetic heart disease is hypertrophic cardiomyopathy (HCM), which 15 is caused by variants in cardiac sarcomeric genes and leads to abnormal heart muscle thickening. 16 Complications of HCM include heart failure, arrhythmia, and sudden cardiac death. The dominant-17 negative c.1208 G>A (p.R403Q) pathogenic variant (PV) in β -myosin (MYH7) is a common and 18 well-studied PV that leads to increased cardiac contractility and HCM onset. Here we identify an 19 20 adenine base editor (ABE) and single-guide RNA system that can efficiently correct this human 21 PV with minimal bystander editing and off-target editing at selected sites. We show that delivery 22 of base editing components rescues pathological manifestations of HCM in induced pluripotent stem cell (iPSC)-cardiomyocytes derived from HCM patients and in a humanized mouse model of 23 24 HCM. Our findings demonstrate the use of base editing to treat inherited cardiac diseases and 25 prompt the further development of ABE-based therapies to correct a variety of monogenic variants causing cardiac disease. 26

28 Main

Hypertrophic cardiomyopathy (HCM), a disease of abnormal heart muscle thickening, is the most 29 common form of genetic heart disease in the United States, affecting upwards of 1 in 200 people 30 ^{1,2}. Clinical complications of HCM include heart failure, arrhythmia, and sudden cardiac death. 31 There is no cure for HCM, aside from heart transplant, which presents its own complications and 32 33 requires lifetime immunosuppression. While HCM-causing variants are found in various sarcomeric protein-encoding genes, over one-third of all HCM-causing variants occur in the 34 Myosin Heavy Chain 7 (*MYH7*) gene, which encodes $\frac{1}{100}$ - β -myosin heavy chain, a motor ATPase 35 36 that incorporates into the thick filament of cardiac muscle and plays a major role in cardiac contraction. These pathogenic variants are generally autosomal dominant missense variants, which 37 allow the incorporation of pathogenic myosin heads into cardiac sarcomeres and lead to increased 38 39 energy consumption, hypercontractility, and disease progression of HCM³.

The heterozygous MYH7 c.1208 G>A (p.R403Q) pathogenic missense variant causes 40 severe HCM with early-onset and progressive myocardial dysfunction and has a high incidence of 41 early sudden cardiac death as fifty percent of patients die by 40 years of age ^{4,5}. It was the first 42 43 MYH7 variant linked to HCM, leading to the discovery of multiple other missense variants in cardiac sarcomere genes and generating numerous studies of its function. The R403Q missense 44 variant, located on the myosin mesa of the myosin head, results in a loss of positive charge on the 45 myosin head, weakening its interaction with myosin binding protein-C, a molecular brake, and 46 leading to an increase in the number of functionally accessible myosin heads available for 47 contraction that pathologically augments sarcomere contractility ⁶. This poison peptide effect 48 results in a dominant-negative disease that requires direct correction or ablation of the pathogenic 49 allele. 50

Base editing has emerged as an attractive method to correct and potentially cure genetically based diseases, especially single nucleotide variants. Base editors are fusion proteins of Cas9 nickase or deactivated Cas9 and a deaminase protein, which allow base pair edits without doublestrand breaks within a defined editing window in relation to the protospacer adjacent motif (PAM) site of a single-guide RNA (sgRNA) ^{7,8}. Adenine base editors (ABEs) use deoxyadenosine deaminase to convert DNA A•T base pairs to G•C base pairs via an inosine intermediate and have been previously shown to function in many post-mitotic cells in vivo and in vitro ⁹⁻¹¹.

Here, we and colleagues who authored the accompanying manuscript (Reichart, Newby, Wakimoto, et al.)¹² report the use of different gene editing strategies to correct the *MYH7* c.1208 G>A (p.R403Q) pathogenic missense variant. We develop an ABE-mediated strategy that efficiently corrects and rescues pathological phenotypes of HCM in patient-derived cells. Furthermore, we generate a humanized mouse model containing the pathogenic missense variant with sequence complementarity to the human sgRNA and show that postnatal ABE correction of this mouse model prevents HCM onset.

65

66 **Results**

67 Identification of an ABE to correct the R403Q pathogenic variant in human iPSCs

To screen various ABEs for their efficiencies, we first inserted the *MYH7* c.1208 G>A (p.R403Q) pathogenic missense variant using CRISPR-Cas9-based homology-directed repair in a human induced pluripotent stem cell (iPSC) line derived from a healthy donor (HD^{*WT*}). We isolated an isogenic heterozygous clone (HD^{403/+}) that mirrors the heterozygous genotype found in patients ⁴, as well as an isogenic homozygous clone (HD^{403/403}) that has not been previously described in patients. Sequencing confirmed no mutations on the highly homologous *MYH6* gene during

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generation of these clones, and these iPSCs readily differentiate into cardiomyocytes (CMs) (Extended Data Fig. 1).

As ABEs have an optimal activity window in protospacer positions 14-17 (counting the 76 first nucleotide immediately 5' of the PAM sequence as protospacer position 1), we chose a sgRNA 77 with an NGA PAM that places the MYH7 c.1208 G>A pathogenic variant in protospacer position 78 79 16 (h403 sgRNA) (Fig. 1a). To identify an optimal ABE capable of efficiently correcting the pathogenic nucleotide back to the wildtype nucleotide without introducing any bystander edits, we 80 tested various engineered deaminases including either ABEmax¹³, which is an optimized, narrow-81 windowed ABE7.10 variant, or ABE8e¹⁴, which is a highly processive, wide-windowed, evolved 82 ABE7.10 variant. Each engineered deaminase variant was fused to engineered SpCas9 variants 83 including SpRY, which targets NRN PAMs ¹⁵; SpG, which targets NGN PAMs ¹⁵; SpCas9-NG, 84 which targets NG PAMs ¹⁶; or SpCas9-VRQR, which targets NGA PAMs ¹⁷. We then screened 85 these ABEs for their efficiency of correction in our HD^{403/403} iPSC line via transient transfection 86 with h403 sgRNA (Fig. 1b). We selected the HD^{403/403} iPSC line for screening to ensure that all 87 G nucleotide Sanger sequencing reads at position c.1208 are due to ABE editing. Similar editing 88 efficiency of the pathogenic adenine was achieved with all ABEmax-SpCas9 variants tested, 89 90 ranging from $26 \pm 2.3\%$ with ABEmax-SpRY to $34 \pm 2.5\%$ with ABEmax-VRQR, with minimal by stander editing of neighboring adenines (the average across three by standers was $2.6 \pm 1.7\%$). 91 92 ABE8e-SpCas9 variants achieved higher editing efficiencies, ranging from $27 \pm 2.6\%$ with 93 ABE8e-SpRY to $37 \pm 1.5\%$ with ABE8e-SpG, with slightly increased by stander editing of neighboring adenines (the average across three bystanders was $4.0 \pm 2.0\%$) (Fig. 1c). These 94 bystander edits are predicted to result in K405E, K405R, or K405G variants in β-myosin heavy 95 chain depending on the combination of edits, which may be deleterious due to the high intolerance 96

of *MYH7* for missense variants ¹⁸, although the consequences of these variants on β -myosin heavy chain function have not been described. For subsequent experiments, we opted to use the more narrow-windowed ABEmax to reduce potential bystander edits, and the SpCas9-VRQR variant with its more stringent PAM requirements to reduce potential Cas-dependent off-target editing.

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102 Correction efficiency and off-target DNA editing analysis in HCM patient-derived iPSCs.

103 To apply our ABEmax-VRQR and h403 sgRNA system to a disease model, we derived human induced pluripotent stem cells (iPSCs) from two HCM patients with the MYH7^{403/+} pathogenic 104 variant (HCM1^{403/+} and HCM2^{403/+}) and corrected the *MYH7*^{403/+} variant via plasmid nucleofection</sup>105 of ABEmax-VRQR-P2a-EGFP and h403 sgRNA, and fluorescence-activated cell sorting of GFP⁺ 106 107 cells (Fig. 2a). By high throughput sequencing (HTS), despite 98-99% on-target editing, we observed minimal to no off-target DNA editing (0.12% or less) at all 58 adenine bases for eight 108 tested candidate off-target loci, which were identified using the bioinformatic tool CRISPOR¹⁹ 109 (Fig. 2b, and Extended Data Fig. 2). A low frequency (0.03-0.48%) of bystander editing was 110 observed at the three bystander adenines for amino acid 505 (K505) of β-myosin. We isolated 111 corrected clonal lines of the HCM patient-derived iPSCs (HCM1^{WT} and HCM2^{WT}) containing no 112 bystander edits or editing of the highly homologous MYH6 gene, ensuring that subsequent 113 characterizations are due to correction of the pathogenic nucleotide. These results suggest that 114 h403 sgRNA with ABEmax-VRQR can efficiently and specifically correct the target pathogenic 115 116 missense variant with minimal bystander editing and little to no DNA editing at tested off-target sites. 117

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119 Functional analyses of ABE-corrected patient-derived iPSC-CMs

In individual CMs, the *MYH7* p.R403Q pathogenic variant increases the number of functional myosin heads available for contraction, which leads to increased force generation and greater ATP consumption. Increases in both force generation and energy consumption have been previously shown in isogenic iPSC-CMs engineered to have the R403Q pathogenic variant from a WT line $^{20-22}$. To determine if these previously validated changes are rescued in our corrected HCM patientderived iPSC-CMs, we differentiated both *MYH7*^{403/+} pathogenic and *MYH7*^{WT} healthy clonal lines for all three patient-derived lines (HD, HCM1, and HCM2) into CMs (**Fig. 2a**).

To investigate whether gene editing correction could reduce hypercontractile force 127 generation in our HCM patient-derived lines, we plated iPSC-CMs at single-cell density on soft 128 polydimethylsiloxane surfaces, recorded high frame-rate videos of contracting CMs, and 129 calculated peak systolic force. The HD^{403/+} iPSC-CMs showed a 1.7-fold increase in peak systolic 130 force compared to HD^{WT} iPSC-CMs originally derived from a healthy donor, consistent with 131 reports of a single MYH7 p.R403Q variant being sufficient to lead to hypercontractility ^{6,22}. On the 132 other hand, corrected HCM1^{WT} and HCM2^{WT} CMs showed a 2.0-fold and 1.6-fold decrease in peak 133 systolic force, respectively, compared to their isogenic HCM1^{403/+} and HCM2^{403/+} counterparts. 134 (Fig 2c). 135

As previous studies have shown that the p.R403Q HCM variant leads to increased ATP consumption and altered cellular metabolism ²³, we next assessed changes in cellular energetics via metabolic flux assays following gene editing correction. Basal oxygen consumption rates (OCR) were increased 1.6-fold in HD^{403/+} iPSC-CMs compared to HD^{WT} iPSC-CMs, and HD^{403/+} iPSC-CMs had a 2.1-fold increase in maximum OCR compared to HD^{WT} iPSC-CMs. Corrected HCM1^{WT} and HCM2^{WT} CMs showed a 1.4-fold and 1.2-fold reduction in basal OCR, respectively, and a 3.7-fold and 2.1-fold reduction in maximum OCR, respectively, compared to isogenic HCM1^{403/+} and HCM2^{403/+} CMs (**Fig. 2d**). These data demonstrate that correction of the pathogenic variant in human HCM CMs is sufficient to rescue the hypercontractility phenotype and restore normal cellular energetics.

146

147 Development of a humanized mouse model of HCM

We next sought to apply base editing to a mouse model of HCM. While β -myosin heavy chain is 148 149 the dominant myosin isoform found in adult human hearts, the highly homologous α -myosin heavy chain is the dominant myosin isoform expressed in adult mouse hearts and is encoded by the *Myh6* 150 gene ²⁴. Consequently, previously described mouse models for HCM have placed the 151 corresponding human MYH7 variant on the mouse Mvh6 gene to account for these expression 152 differences. While the 35 amino acids around R403 are 100% identical between human MYH7 and 153 154 mouse Myh6, the DNA sequence encoding this region of the protein is not identical (Extended Data Fig. 3). Thus, sgRNAs and editing strategies developed for the human genome would not be 155 directly applicable to a mouse model. 156

To perform preclinical studies using our human sequence-specific base editing strategy, 157 we generated a humanized mouse model containing the MYH7 c.1208 G>A (p.R403Q) human 158 missense variant within the mouse Myh6 gene that also has human DNA sequence identity of at 159 least 21 nucleotides upstream and downstream from the pathogenic variant to allow testing of 160 human genome specific CRISPR strategies (Fig. 3a). The other Myh6 allele contained the 161 unmodified mouse genomic sequence. This humanized mouse model $(Mvh6^{h403/+})$ mirrors the 162 phenotype of previously described *Myh6* p.R403Q mouse models ^{25,26}. Most notably, homozygous 163 mice $(Myh6^{h403/h403})$ have enlarged atria, extensive interstitial fibrosis, and die within the first week 164 of life (Fig. 3b). At 9 months of age, $Myh6^{h403/+}$ mice have developed cardiomyopathy with 165

166 significant ventricular hypertrophy, myocyte disarray, and fibrosis (**Fig. 3c**).

167

168 In vivo ABE treatment of a mouse model of human HCM

We packaged ABEmax-VRQR and h403 sgRNA within adeno-associated virus (AAV), a US 169 Food and Drug Administration approved viral delivery method. We chose to use the AAV9 capsid 170 due to its high cardiac transduction and use in clinical trials ²⁷. As the full-length base editor (~5.4 171 kb) exceeds the packaging limit of a single AAV9 (\sim 4.7 kb), we split the base editor across two 172 AAV9s and used *trans*-splicing inteins to reconstitute the full-length base editor in cells upon 173 protein expression ²⁸. As AAV9 contains broad tissue tropism, we utilized a cardiac troponin T 174 (cTnT) promoter to limit expression of the base editor to CMs ²⁹. For this dual AAV9 system, each 175 AAV9 also contained a single copy of an expression cassette encoding h403 sgRNA (Fig. 4a). 176

We first sought to validate the efficiency of our dual AAV9 ABE system by trying to rescue 177 $Myh6^{h403/h403}$ mice, which die within the first week of life. Notably, no human patients have been 178 reported to have the homozygous genotype. We injected P0 (postnatal day 0) *Myh6*^{h403/h403} pups 179 intrathoracically with either saline, a low dose $(4 \times 10^{13} \text{ vg/kg})$, or a high dose $(1.5 \times 10^{14} \text{ vg/kg})$ of 180 each AAV9 (total of 8×10^{13} vg/kg for low, and 3×10^{14} vg/kg for high) and monitored their 181 development (Extended Data Fig. 4a). The 3×10^{14} vg/kg high dose is the highest dose 182 administered in clinical trials ³⁰. The $Myh6^{h403/+}$ and $Myh6^{WT}$ mice survived past weaning and well 183 into adulthood. The median survival of saline-injected $Myh6^{h403/h403}$ mice was 7.0 days, whereas 184 that of low-dose ABE-treated Myh6h403/h403 mice was increased to 9.0 days (1.3-fold longer, 185 P < 0.05 by Mantel-Cox test). The median survival of high-dose ABE-treated $Mvh6^{h403/h403}$ mice 186 was increased to 15.0 days (2.1-fold longer, P < 0.01 by Mantel-Cox test) (Extended Data Fig. 187 4b). Sanger sequencing of cDNA of the heart from a high-dose mouse indicated 35% correction 188

of the pathogenic nucleotide at the transcript level, suggesting that our dual AAV9 ABE system
enabled editing in the heart (Extended Data Fig. 4).

As the MYH7 p.R403Q variant only exists in a heterozygous form in human patients, we 191 next deployed our dual AAV9 ABE system to prevent HCM disease onset in $Myh6^{h403/+}$ mice. We 192 injected $Myh6^{h403/+}$ P0 pups intrathoracically with either saline or 1×10^{14} vg/kg of each AAV9 193 $(2 \times 10^{14} \text{ vg/kg total})$ and their littermate $Myh6^{WT}$ control pups with saline. At 5 weeks of age, the 194 mice were put on a chow diet containing 0.1% cyclosporine A, which has previously been shown 195 to accelerate the onset of HCM in mouse models of sarcomere pathogenic variants ³¹. Serial 196 echocardiograms were conducted at 8, 12, and 16 weeks of age to monitor disease progression 197 (Fig. 4b). We found that $Myh6^{h403/+}$ mice had increased features of HCM compared to $Myh6^{WT}$ 198 199 controls, including increased left ventricular anterior wall thickness at diastole (LVAW;d) (1.07 \pm 0.0443 mm vs. 0.883 \pm 0.0441 mm, P = 0.017) and increased left ventricular posterior wall 200 thickness at diastole (LVPW;d) (1.04 \pm 0.0809 mm vs. 0.867 \pm 0.0590 mm, P = 0.128). These 201 202 mice also had decreased left ventricular internal diameter at diastole (LVID;d) (2.34 ± 0.142 mm vs. 2.81 ± 0.0540 mm, P = 0.015) and systole (LVID;s) (0.940 ± 0.0713 mm vs. 1.24 ± 0.0520 , P 203 204 = 0.010), with slightly increased ejection fraction (EF) and fractional shortening (FS). The increased ventricular wall thickness and a concomitant decrease in ventricular diameter of 205 $Myh\delta^{h403/+}$ mice, along with high-normal fractional shortening is consistent with the clinical 206 progression in human patients ¹. 207

In contrast, ABE-treated $Myh6^{h403/+}$ mice had reduced features of HCM with comparable similar echocardiographic measurements to $Myh6^{WT}$ control mice, suggesting that gene correction of the pathogenic nucleotide was sufficient to prevent the onset of HCM (**Fig. 4c-h**, **Table 1**, and **Extended Data Fig. 5**). Histological analysis also revealed increased cardiac wall thickness and

decreased ventricular diameter cross-sectional area in $Myh6^{h403/+}$ mice compared to $Myh6^{WT}$ 212 control mice, while ABE-treated $Myh6^{h403/+}$ mice had similar cardiac dimensions to $Myh6^{WT}$ 213 control mice (Fig. 4i-k). When normalized to tibia length, $Myh6^{h403/+}$ mice had 1.3-fold larger 214 hearts by heart weight compared to $Myh6^{WT}$ control mice, while ABE-treated $Myh6^{h403/+}$ mice had 215 no significant difference in heart weight compared to $Myh6^{WT}$ mice (Fig. 41). As a measure of 216 fibrosis, hearts from $Myh6^{h403/+}$ mice had 3.0-fold more collagen area compared to $Myh6^{WT}$ control 217 mice, while ABE-treated $Myh6^{h403/+}$ mice had no significant difference in collagen area compared 218 to *Mvh6^{WT}* mice (Fig. 4m). These data suggest that dual AAV9 ABE treatment was sufficient to 219 prevent the onset of HCM-mediated pathological remodeling of the heart. 220

221

222 Genomic and transcriptomic analyses of ABE-treated mice.

To determine tissue-level viral transduction and gene editing efficiency, we collected the heart, 223 lung, liver, spleen, and quadriceps muscle from saline-treated $Myh6^{h403/+}$ mice, and ABE-treated 224 $Myh6^{h403/+}$ mice at 18 weeks of age. The heart was further dissected into the right atrium, right 225 ventricle, left atrium, and left ventricle, and each chamber was separately analyzed. We conducted 226 a viral copy number assay and found similar viral transduction in all four chambers of the heart, 227 228 ranging from 5-9 copies of each viral half per diploid genome. The liver had the highest viral transduction of the dissected tissues, with about 10 copies of each viral half per diploid genome, 229 230 while the spleen had the least, with about 0.2 copies of each viral half per diploid genome 231 (Extended Data Fig. 6a). Despite similar levels of viral copy numbers among tissues, we found low levels of DNA editing in the liver, lung, spleen, and quadriceps (<0.12%) compared to DNA 232 editing in each of the heart chambers (5.5-8.0%), confirming the cardiac specificity of the cTnT 233 promoter (Extended Data Fig. 6b). As CMs make up only ~30% of cells ³² in the heart but are 234

the only cells to express Myh6, we next looked at cDNA efficiency. We found that while the left 235 atria had less correction of pathogenic transcripts compared to the ventricles, there was similar 236 editing efficiency across the four chambers (12.9-26.7%) (Extended Data Fig. 6c). We further 237 confirmed no changes in protein expression of key sarcomere proteins (Extended Data Fig. 6d). 238 To identify genomic and transcriptomic changes more deeply in CMs following base 239 editing, we isolated CM nuclei from the ventricles of saline-treated Myh6^{WT} control mice, saline-240 treated $Myh6^{h403/+}$ mice, and ABE-treated $Myh6^{h403/+}$ mice at 18 weeks of age (Fig. 5a). We first 241 evaluated on-target editing efficiencies following dual AAV9 ABE treatment. In ABE-treated 242 $Mvh6^{h403/+}$ mice, DNA editing efficiency of the target pathogenic adenine was $32.3 \pm 2.87\%$, 243 resulting in a 33.1 \pm 9.08% reduction in pathogenic transcripts compared to *Myh6*^{h403/+} mice (Fig. 244 **5b,c**), which is comparable to other *in vivo* studies using base editing ²⁶ or RNAi-based knockdown 245 of pathogenic transcripts ³³. Furthermore, there was no detectable bystander editing in ABE-treated 246 $Myh6^{h403/+}$ mice (Fig. 5d). We then assessed potential off-target RNA editing via transcriptome-247 248 wide RNA sequencing (RNA-seq), as ABEmax contains deaminase activity. RNA-seq analysis revealed no significant increase in the average frequency of A-to-I editing in the transcriptome of 249 ABE-treated mice compared to that of saline-treated mice (Fig. 5e). This finding suggests that in 250 251 vivo treatment with our dual AAV9 ABE system does not increase RNA deamination above background levels of endogenous cellular deaminase activity. 252

We next evaluated transcriptome-wide changes in ABE-treated $Myh6^{h403/+}$ mice via RNAseq. We first identified 257 differentially regulated genes between $Myh6^{WT}$ mice and $Myh6^{h403/+}$ mice. Heat maps showed that ABE-treated $Myh6^{h403/+}$ mice had transcriptome profiles more similar to $Myh6^{WT}$ mice than to $Myh6^{h403/+}$ mice in a correction efficiency-dependent response (Fig. 5f, and Extended Data Fig. 7). Gene ontology analyses of differentially regulated genes between 258 $Myh6^{h403/+}$ mice and $Myh6^{WT}$ mice indicated dysregulation of intercellular signaling and 259 angiogenesis, while intercellular signaling was dysregulated between $Myh6^{h403/+}$ mice and ABE-260 treated $Myh6^{h403/+}$ mice (**Fig. 5g**). Additionally, expression of the prototypic hypertrophic marker 261 Nppa was 2.8-fold higher in $Myh6^{h403/+}$ mice compared to $Myh6^{WT}$ mice, while expression of Nppa262 in the ABE-treated $Myh6^{h403/+}$ mice was not significantly different from $Myh6^{WT}$ mice (**Fig. 5h**). 263 Taken together, these data suggest that our dual AAV9 ABE system can efficiently correct the 264 pathogenic nucleotide in genomic DNA and prevent transcriptomic dysregulation.

265

266 **Discussion**

In this proof-of-concept study, we show that an adenine base editor can directly correct a common 267 and well-studied pathogenic variant that leads to HCM, serving as a first step toward clinical 268 translation of base editing in HCM patients. In iPSCs derived from patients with clinically 269 diagnosed HCM, we identified the ABEmax-VRQR base editor as the optimal base editor, as it 270 corrected the target pathogenic adenine at 98-99% efficiency in iPSCs sorted to contain all editing 271 components with minimal bystander editing of neighboring adenines and low DNA off-target 272 editing at tested sites within the human genome. These corrected patient-derived iPSCs, when 273 274 differentiated into CMs, demonstrated normalization of contractile force, and rescued cellular energetics compared to healthy controls uncorrected patient-derived CMs, suggesting that 275 correction of this pathogenic variant is sufficient to reduce the abnormalities of HCM. 276

To extend this work to a clinically relevant animal model, we generated a humanized mouse model of HCM. This humanized mouse model demonstrates similar HCM onset to other previously described models, but contains DNA sequence complementarity to the human allele, thus allowing testing of human-specific sgRNAs. Injection of a single dose of dual AAV9 encoding for ABEmax-VRQR and the h403_sgRNA in clinically relevant heterozygous postnatal $Myh6^{h403/+}$ mice resulted in correction of the pathogenic allele, reduction of pathogenic transcripts, and reduction in cardiac hypertrophy and histopathologic remodeling. Our study, along with the study from our colleagues ¹², marks the first demonstration of efficient single nucleotide gene correction in vivo in postnatal mammalian cardiomyocytes. Future studies will seek to determine whether this rescue is sufficient to prevent HCM onset for the lifespan of the mouse.

Base editing in the $Myh6^{h403/h403}$ homozygous mice could double their lifespan, but 287 ultimately could not prevent their death before weaning. This is predictable, since only the 288 heterozygous form of the MYH7 p.R403Q variant exists in human patients, and most likely, the 289 homozygous form is nonviable. In mice, Myh7 protein is partially expressed within the ventricles 290 for the first week of life, which likely protects the heart against the pathogenic Myh6 protein ²⁴, as 291 these mice die around 7 days after birth coincident with isoform switching in the ventricles. While 292 we could achieve ~35% correction of transcripts at the highest dual AAV9 dose, the lag between 293 healthy protein production following correction and myosin turnover and replacement of the 294 pathogenic protein was likely too long to prevent death. 295

As HCM is an autosomal dominant disease, there is a strong founder effect bias in patient 296 297 frequency; the initial study linking the c.1208G>A pathogenic variant to HCM was based on genetic mapping of a large French-Canadian kindred (>80 members) over five generations ³⁴. 298 299 Nonetheless, this pathogenic variant is likely found worldwide as it was also reported in a Korean family ³⁵. Based on data from the Sarcomeric Human Cardiomyopathy Registry (SHaRe) ³⁶, which 300 performed genetic testing on HCM patients across eight different institutions in three different 301 302 continents, out of all patients with identified MYH7 gene variants, 2.4% (15/613) of them had the 303 c.1208G>A pathogenic variant, making it the fifth most frequent pathogenic variant out of 229

MYH7 variants detected. For all sarcomere gene variants identified, 0.5% (15/2763) of patients 304 had the c.1208G>A pathogenic variant. The World Bank estimates that there are 5.1 billion people 305 306 worldwide aged 15-64 (https://databank.worldbank.org/source/population-estimates-andprojections). It is conservatively estimated that 1 in 500 people have HCM, of which 50% have a 307 known genetic cause in a sarcomere gene. From these data, we can estimate that perhaps 25,000 308 309 patients exist with the specific c.1208G>A p.R403Q pathogenic variant and would be amenable to adenine base editing correction. 310

Although we demonstrate base editing for a single pathogenic missense variant in MYH7, 311 we envision that this base editing approach can be applied to the hundreds of other documented 312 missense variants in MYH7 and to other cardiac sarcomeric proteins in which dominant-negative 313 variants lead to HCM ¹⁸. Base editing is an ideal tool to correct the many dominant-negative 314 cardiac diseases, as gene therapy is precluded in dominant-negative diseases, and CRISPR-Cas9 315 single-cutting may edit the healthy wildtype allele, which can have deleterious effects for essential 316 genes such as MYH7³⁷. In base editing, careful selection of a sgRNA and a base editor with 317 appropriate editing window and activity can minimize off-target binding of the sgRNA to the 318 healthy wildtype allele and minimize DNA off-target and bystander editing. As a clinical 319 320 treatment, we envision that while the initial base editing trials in the heart will be rigorously evaluated before clinical translation, these and other initial studies will establish a pipeline for gene 321 322 editing of HCM. In the future, treatment of other MYH7 pathogenic variants may then simply 323 involve changing the Cas9 variant and gRNA sequence and evaluating potential gRNA-specific off-target editing for each patient's genome. 324

While our study showed that gene editing of newborn mice can prevent the onset of HCM caused by the highly penetrant p.R403Q variant, some patients are genotype-positive for an HCM-

causing variant but are phenotype-negative by not demonstrating LV hypertrophy at any point in 327 the clinical course ³⁸. These patients would be unlikely to need gene editing correction unless 328 clinical HCM develops. Future studies will seek to determine whether in vivo gene editing in 329 established HCM can either revert or limit the progression of the disease. Previous studies in a 330 mouse ³⁹ and cat model ⁴⁰ of HCM have shown that mavacamten, a small molecule myosin 331 inhibitor, can cause partial regression of hypertrophy in adult animals with established HCM, 332 suggesting a potentially wide therapeutic window for other treatments such as gene editing. Use 333 of adult animals with larger hearts could also allow testing of other delivery methods not explored 334 in our study, such as coronary injection ⁴¹, which could then allow testing of other delivery 335 modalities for base editors including lipid nanoparticles ⁴² or virus-like particles ⁴³. Our study also 336 did not explore optimization in viral dosage, use of a non-targeting sgRNA as a control treatment 337 in place of saline administration, and or use of other cardiogenic AAV capsid variants ^{44,45}. Finally, 338 future studies will need to demonstrate correction in larger animal models that endogenously 339 express β -myosin heavy chain predominantly within the heart, such as rabbits, cats, $\frac{\theta}{\theta}$ and pigs $\frac{46}{2}$, 340 before ultimately treating HCM patients. Myosin-binding drugs that can modulate cardiac 341 function, such as mavacamten ³⁹, hold great promise in treating HCM; clinical trials for 342 mavacamten have met their primary endpoints, and it was recently approved by the FDA to treat 343 symptomatic obstructive HCM ^{47,48}. Nevertheless, there is a continued need for new therapeutic 344 modalities, especially considering the on-label warning of heart failure of mavacamten ⁴⁹. Our 345 study suggests that base editing can provide an alternative one-time treatment modality to directly 346 and permanently correct pathogenic HCM-causing variants and prevent HCM onset, especially in 347 cardiac genes for which no drug has been identified, or for patients with symptomatic non-348

obstructive HCM, for which mavacamten did not significantly improve exercise capacity or
 symptoms compared to placebo ⁵⁰.

351

352 Methods

353 Study design and approval

The objective of this study was to determine whether base editing correction of a pathogenic HCM-354 causing variant could prevent the onset of HCM pathological features in human CMs and a 355 humanized mouse model. In human CMs, this was done by base editing correction of HCM 356 patient-derived iPSCs and measuring changes in characteristic CM function. In a humanized 357 mouse model, a dual AAV9 system was used to deliver the base editing components to CMs and 358 changes in heart function, dimensions, and transcriptomics were measured. For all experiments, 359 the number of replicates, type of replicates, and statistical test used is are reported in the figure 360 legends. For in vitro CM experiments, data are collected from three separate differentiations, and 361 no outliers or other data points were excluded. For in vivo experiments, male and female mice 362 were assigned to treatment based on genotype. Echocardiographic measurements were conducted 363 in a blinded fashion. Runt mice with reduced body weights more than 2 standard deviations from 364 the mean were excluded. Endpoints were guided by changes in echocardiographic measurements. 365 Animal work described in this manuscript has been approved and conducted under the oversight 366 of the UT Southwestern Institutional Animal Care and Use Committee. 367

368

369 **Plasmids and vector construction**

The pSpCas9(BB)-2A-GFP (PX458) plasmid was a gift from Feng Zhang (Addgene plasmid #48138)⁵¹, and was used as the primary scaffold to clone in the following base editors and SpCas9

nickases: ABE8e, a gift from David Liu (Addgene plasmid #138489)¹⁴; VRQR-ABEmax, a gift 372 from David Liu (Addgene plasmid #119811)⁵²; NG-ABEmax, a gift from David Liu (Addgene 373 plasmid #124163) ⁵²; pCMV-T7-SpG-HF1-P2A-EGFP (RTW5000), a gift from Benjamin 374 Kleinstiver (Addgene plasmid #139996)¹⁵; and pCMV-T7-SpRY-HF1-P2A-EGFP (RTW5008), 375 a gift from Benjamin Kleinstiver (Addgene plasmid #139997)¹⁵. The N-terminal ABE and C-376 377 terminal ABE constructs were adapted from Cbh v5 AAV-ABE N terminus (Addgene plasmid #137177) ⁵³ and Cbh v5 AAV-ABE C terminus (Addgene plasmid #137178) ⁵³ and synthesized 378 by Twist Bioscience. PCR amplification of select plasmids was done using PrimeStar GXL 379 Polymerase (Takara), and cloning was done using NEBuilder HiFi DNA Assembly (NEB) into 380 restriction enzyme-digested destination vectors. 381

382

383 Generation of patient-derived iPSCs and isogenic lines

Peripheral blood mononuclear cells (PBMCs) from two patients with the MYH7 c.1208 G>A 384 (p.R403Q) pathogenic variant were reprogrammed to iPSCs (HCM1 and HCM1HCM2) using 385 Sendai virus. The HCM1 line was derived from a 56-year-old female with extensive family history 386 of HCM and nonobstructive HCM with a history of reduced left ventricular ejection fraction and 387 388 low maximal oxygen uptake (VO_2 max). A biventricular pacemaker was placed for a complete heart block. The HCM2 line was derived from a 32-year-old male with a history of HCM, an 389 390 implantable cardioverter-defibrillator, and a strong family history of HCM. He has a dilated left 391 atrium but has improved VO₂ max, metabolic equivalent (METs), and no evidence of atrial fibrillation by cardiopulmonary exercise testing. These two human iPSC lines were obtained from 392 393 Joseph C. Wu, MD, PhD at the Stanford Cardiovascular Institute funded by NIH R24 HL117756. 394 PBMCs from a healthy male donor (HD) were reprogrammed to iPSCs at the UT Southwestern

Wellstone Myoediting Core using Sendai virus (CytoTune 2.0 Sendai Reprogramming Kit, 395 ThermoFisher Scientific). To generate isogenic iPSCs containing the MYH7 c.1208 G>A 396 (p.R403Q) variant via homology-directed repair, HD iPSCs were nucleofected using the P3 397 Primary Cell 4D-Nucleofector X Kit (Lonza) with a single-stranded oligodeoxynucleotide 398 399 (ssODN) template (Integrated DNA Technologies, IDT) encoding for the pathogenic variant, and 400 the PX458 plasmid encoding SpCas9-P2a-EGFP and a sgRNA targeting MYH7. For base editing correction of HCM1 and HCM2 patient derived lines, iPSCs were nucleofected with a single 401 402 plasmid encoding for ABEmax-VRQR-P2a-EGFP and h403 sgRNA. After 48 hours, GFP+ iPSCs were collected by fluorescence-activated cell sorting, clonally expanded, and genotyped by Sanger 403 sequencing. 404

405

406 **iPSC maintenance and differentiation**

iPSC culture and differentiation were performed as previously described ¹¹. Briefly, iPSCs were 407 408 cultured on Matrigel (Corning)-coated tissue culture polystyrene plates and maintained in mTeSR1 media (STEMCELL) and passaged at 70-80% confluency using Versene. iPSCs were 409 differentiated into CMs at 70-80% confluency by treatment with CHIR99021 (Selleckchem) in 410 411 RPMI supplemented with ascorbic acid (50 µg/mL) and B27 without insulin (RPMI/B27-) for 24 hrs (from day (d) 0 to d1). At d1, media was replaced with RPMI/B27-. At d3, cells were treated 412 413 with RPMI/B27- supplemented with WNT-C59 (Selleckchem). At d5, media was refreshed with 414 RPMI/B27-. From d7 onwards, iPSC-CMs were maintained in RPMI supplemented with ascorbic acid (50 µg/mL) and B27 (RPMI/B27) with media refreshed every 3-4 days. Metabolic selection 415 416 of CMs was performed for 6 days starting d10 by culturing cells in RPMI without glucose and 417 supplemented with 5 mM sodium DL-lactate and CDM3 supplement (500 µg/mL Oryza sativaderived recombinant human albumin, A0237, Sigma-Aldrich; and 213 μ g/mL L-ascorbic acid 2phosphate, Sigma-Aldrich). To induce their maturation, iPSC-CMs were maintained in RPMI without glucose supplemented with B27, 50 μ mol palmitic acid, 100 μ mol oleic acid, 10 mmol galactose, and 1 mmol glutamine (Sigma-Aldrich) ^{54,55}. All CM functional studies were done at d40-50.

423

424 Plasmid transfection and editing efficiency analysis

iPSCs were seeded on a 48-well plate 24 h before transfection. At ~20% confluency, cells were transiently transfected with 0.5 μ g of plasmid encoding for a base editor and the h403_sgRNA using 1 μ L of Lipofectamine Stem Transfection Reagent (ThermoFisher) per well. Following 48 h post-transfection, cells were lysed in Direct PCR Lysis Reagent (Cell) (Viagen). PCR amplification of target sites was done using PrimeStar GXL Polymerase (Takara), and PCR cleanup was done using ExoSap-IT Express (ThermoFisher) before Sanger sequencing. Chromatograms were analyzed using EditR to determine base editing efficiencies ⁵⁶.

432

433 Contractility analyses of iPSC-CMs

iPSC-CMs were plated at single-cell density on flexible polydimethylsiloxane (PDMS) 527
substrates (Young's modulus = 5 kPa) prepared according to a previously established protocol ⁵⁷.
Recordings of contracting iPSC-CMs were captured at 37 °C using a Nikon A1R+ confocal system
at 59 frames per second in resonance scanning mode. Contractile force generation of iPSC-CMs
was quantified using a previously established method. In brief, recordings were analyzed using
Fiji to measure maximum and minimum cell lengths, and cell widths during contraction. A
previously published customized MATLAB code was used to calculate peak systolic forces ⁵⁸.

442 Extracellular flux analyses of iPSC-CMs

iPSC-CMs were plated at 40,000 cells per well in Seahorse XFe96 V3 PS Cell Culture Microplates 443 (Agilent) coated with Matrigel. One-week post-plating, cells were washed three times with 444 prewarmed assay media (pyruvate-free DMEM (Sigma D5030) supplemented with 2 mM L-445 446 glutamine, 1 mM sodium pyruvate, and 10 mM glucose, pH 7.4) and incubated at 37 °C for 60 min in a non-CO₂ incubator. Oxygen consumption rate (OCR) was measured in a Seahorse XFe96 447 instrument using consecutive cycles of 2 mins of measurement, 10 seconds of waiting, and 3 448 minutes of mixing. Mitochondrial stress testing was performed by injecting oligomycin (final 449 concentration 2 μ M), CCCP (final concentration 1 μ M), and antimycin A (final concentration 1 450 µM) at indicated time intervals. Data were analyzed using the WAVE software (Agilent). 451

452

453 Immunofluorescence staining

454 iPSC-CMs were plated on glass surfaces and fixed with 4% paraformaldehyde for 10 min, 455 followed by blocking with 5% goat serum/0.1% Tween-20 (Sigma-Aldrich) for 1 hr. Primary and 456 secondary antibodies were diluted in blocking buffer and added to cells for 2 hr and 1 hr, 457 respectively. Nuclei were counterstained using DAPI. Antibodies used included rabbit anti-458 troponin-I (H-170 sc-15368, Santa Cruz Biotechnology, 1:200), and fluorescein-conjugated 459 donkey anti-rabbit IgG (711-095-152, Jackson ImmunoResearch, 1:50).

460

461 **Off-target analyses**

462 Candidate off-target sites were identified with CRISPOR, and the top 8 sites by cutting frequency
 463 determination (CFD) score, for which PCR products were successfully obtained, were selected

^{19,59}. Genomic DNA was isolated using a DNeasy Blood & Tissue Kit (Qiagen) from HCM1 and 464 HCM2 cell lines that had been nucleofected with plasmids encoding for ABEmax- VRQR-P2a-465 EGFP and h403 sgRNA and sorted for GFP+ cells. Target sites were PCR amplified using 466 PrimeStar GXL Polymerase (Takara), and a second round of PCR was used to add Illumina flow 467 cell binding sequences and barcodes. PCR products were purified with AMPure XP Beads 468 469 (Beckman Coulter), analyzed for integrity on a 2200 TapeStation System (Agilent), and quantified by QuBit dsDNA high-sensitivity assay (Invitrogen) before pooling and loading onto an Illumina 470 MiSeq. Following demultiplexing, resulting reads were analyzed with CRISPResso2 for editing 471 frequency ⁶⁰. 472

473

474 Generation of adeno-associated viruses

Recombinant AAV9 (rAAV9) viruses were made at the University of Michigan Vector Core using
ultracentrifugation through an iodixanol gradient. rAAV9s were washed 3 times with PBS using
Amicon Ultra Centrifugal Filter Units (Millipore) and resuspended in PBS + 0.001% Pluronic F68.
Titers were assessed by qPCR. rAAV9 was stored in 25 µL aliquots at -80 °C.

479

480 Mice

Mice were housed in a barrier facility with a 12-hour:12-hour light:dark cycle and maintained on standard chow (2916 Teklad Global). The humanized $Myh6^{h403/+}$ pathogenic variant was introduced via microinjection of zygotes with Cas9 mRNA (50 ng/µL) (TriLink Biotechnologies), a sgRNA (20 ng/µL) (IDT), and a ssODN donor template (15 ng/µL) (IDT) following a modified protocol ⁶¹. Genotyping was performed using a custom TaqMan SNP Genotyping Assay (Assay ID: ANPRZE6) (ThermoFisher). To accelerate the onset of HCM, mice were treated with a custom chow (2916 Teklad Global base) containing Cyclosporine A (Alfa Aesar) at 1 g/kg and blue food dye at 0.2 g/kg. For injections, mice were genotyped at P0 and received either saline or a AAV9 dose via a single 40 µL bolus using a 31G insulin syringe through the diaphragm by a subxiphoid approach into the inferior mediastinum, avoiding the heart and the lung.

491

492 Transthoracic echocardiography

Cardiac function on conscious mice was evaluated by two-dimensional transthoracic 493 echocardiography using a VisualSonics Vevo2100 imaging system. M-mode tracings were used 494 to measure LV anterior wall thickness at diastole (LVAW;d), LV posterior wall thickness at 495 diastole (LVPW;d), and LV internal diameter at end diastole (LVIDd) and end systole (LVIDs). 496 FS was calculated according to the following formula: FS (%) = $[(LVIDd - LVIDs)/LVIDd] \times$ 497 100. EF was calculated according to the following formula: EF (%) = [(LVEDV -498 LVESV)/LVEDV] x 100. All measurements were performed by an experienced operator blinded 499 500 to the study.

501

502 Histology

Mouse hearts were dissected out and submerged in PBS with cardioplegic 0.2M KCl for 5 minutes before fixation in 4% paraformaldehyde in PBS overnight, followed by dehydration in 70% ethanol and paraffin embedding. Serial transverse cross-sections at 500 µm intervals were cut and mounted on slides, followed by H&E staining, Picrosirius Red, or Masson's Trichrome staining. Images were captured on a BZ-X800 all-in-one microscope (Keyence) at 10x or 40x magnification. Analyses were performed with Fiji and Adobe Photoshop.

509

510 Viral copy number (VCN) assay

Genomic DNA was isolated from mouse tissue using the DNeasy Blood & Tissue Kit (Qiagen). AAV VCN was determined by quantitative PCR (qPCR) using custom-designed primers and Taqman probes (IDT) (**Supplementary Table 1.**) and the Applied Biosystems TaqMan Fast Advanced Master Mix (Applied Biosystems). The primers and probes anneal to the N-terminal and C-terminal Cas9 genes. A copy number standard curve of the AAV plasmids used for packaging was used to determine copy number for each cycle threshold. The 18S ribosomal RNA gene was used as a reference to calculate genomic DNA quantity.

518

519 Cardiac myofibril isolation and analysis

520 Cardiac myofibrils were isolated from hearts using a previously reported protocol, with minor 521 modifications ⁶². Collected hearts were homogenized using CK28-R hard tissue homogenizing 522 tubes in the Percellys homogenizer (Bertin Instruments) on the hard tissue setting. Following 523 myofibril isolation, protein amounts were quantified by Pierce BCA Protein Assay 524 (ThermoFisher). For each sample, 10 μg protein was run on a 4-20% polyacrylamide gel, then 525 stained with Coomassie G-250.

526

527 CM nuclei isolation

For each nuclear sample, ventricular heart tissue was isolated. CM nuclei were isolated as previously described ⁶³. Isolated nuclei were immediately used for downstream processing. RNA was isolated from nuclei using the RNeasy Micro Kit (Qiagen). For qPCR and cDNA HTS, RNA was reverse transcribed using iScript cDNA Synthesis Kit (Bio-Rad). For DNA sequencing, nuclei were lysed in Direct PCR Lysis Reagent (Cell) (Viagen).

534 Mouse DNA and cDNA sequencing and analysis

535 On-target DNA and cDNA sites were PCR amplified using PrimeStar GXL Polymerase (Takara), 536 and a second round of PCR was used to add Illumina flow cell binding sequences and barcodes. 537 PCR products were purified with AMPure XP Beads (Beckman Coulter), analyzed for integrity on 538 a 2200 TapeStation System (Agilent), and quantified by QuBit dsDNA high-sensitivity assay 539 (Invitrogen) before pooling and loading onto an Illumina MiSeq. Following demultiplexing, 540 resulting reads were analyzed with CRISPResso2 for editing frequency ⁶⁰.

541

542 RNA-seq library preparation, sequencing, and analysis

RNA-seq libraries were generated using the SMARTer Stranded Total RNA-Seq Kit v2-Pico Input 543 Mammalian kit (Takara), containing Illumina sequencing adapters. Libraries were visualized on a 544 2200 TapeStation System (Agilent) and quantified by QuBit dsDNA high-sensitivity assay 545 546 (Invitrogen) before pooling and loading onto an Illumina NextSeq 500. FastQC tool (Version 0.11.8) was used for quality control of RNA-seq data to determine low quality or adapteer portions 547 of the reads for trimming. Read trimming was performed using Trimmomatic (Version 0.39) and 548 549 strandness was determined using RSeQC (Version 4.0.0) and then reads were aligned to the mm10 reference genome using HiSAT2 (Version 2.1.0) with default settings and -rna-strandness R. 550 551 Aligned reads were counted using featureCounts (Version 1.6.2). Differential gene expression 552 analysis was performed using R package DESeq (Version 1.38.0). Genes with fold-change >2 and p-value <0.01 were designated as DEGs between sample group comparisons. To calculate the 553 average percentage of A-to-I editing amongst adenosines sequenced in transcriptome-wide 554 sequencing analysis, we adopted a previous strategy ⁹. In brief, REDItools2 was used to quantify 555

the percentage editing in each sample. Nucleotides except adenosines were removed and remaining adenosines with read coverage less than 10 or read quality score below 25 were also filtered to avoid errors due to low sampling or low sequencing quality. We then calculated the number of Ato-I conversion in each sample and divided this by the total number of adenosines in our dataset after filtering to get the percentage of A-to-I editing in the transcriptome.

561

562 **Quantitative real-time PCR analysis**

qPCR reactions were assembled using Applied Biosystems TaqMan Fast Advanced Master Mix
 (Applied Biosystems). Assays were performed using Applied Biosystems QuantStudio 5 Real Time PCR System (Applied Biosystems). Expression values were normalized to *18S* mRNA and
 represented as fold change.

567

568 Statistics

All data are presented as means \pm s.e.m. or means \pm s.d. as indicated. Unpaired two-tailed Student's t tests were performed for comparison between the respective two groups as indicated in the figures. Kaplan-Meier analysis and Log-rank (Mantel-Cox) test were used to evaluate the difference in survival between different genotypes. Data analyses were performed with statistical software (GraphPad Prism Software). *P* values less than 0.05 were considered statistically significant.

575

576 **Data and materials availability:** All data needed to evaluate the conclusions in the paper are 577 present in the paper, extended data, and supplementary materials. Raw and analyzed RNA-578 sequencing data generated during this study are available in the Gene Expression Omnibus

| 579 | (GEO) | repository (http://www.ncbi.nlm.nih.gov/geo/) and are accessible through GEO series | | | | |
|------------|-----------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|--|--|
| 580 | accession number GSE201755. | | | | | |
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| 581 | | | | | | |
| 582 | Code a | vailability: The MATLAB code used to perform contractile force measurements of | | | | |
| 583 | iPSC-C | Ms has been deposited to GitHub: https://github.com/DarisaLLC/Cardio. | | | | |
| 591 | | | | | | |
| 564 | | | | | | |
| 585 | Extend | ed Data | | | | |
| 586 | Extende | ed Data Fig. 1-7. | | | | |
| 587 | Supple | mentary Information | | | | |
| 588 | Suppler | nentary Fig. 1-2. | | | | |
| 589 | Supplementary Table 1. | | | | | |
| 590 | Referei | nces and Notes | | | | |
| 591 | 1. | Maron, B.J. Clinical Course and Management of Hypertrophic Cardiomyopathy. <i>N Engl</i> | | | | |
| 592 593 | 2. | J Med 379, 655-668 (2018). Semsarian, C., Ingles, J., Maron, M.S. & Maron, B.J. New perspectives on the prevalence | | | | |
| 594 | | of hypertrophic cardiomyopathy. J Am Coll Cardiol 65, 1249-1254 (2015). | | | | |
| 595 | 3. | Trivedi, D.V., Adhikari, A.S., Sarkar, S.S., Ruppel, K.M. & Spudich, J.A. Hypertrophic | | | | |
| 596 | | cardiomyopathy and the myosin mesa: viewing an old disease in a new light. <i>Biophys Rev</i> | | | | |
| 597 | 1 | 10, 27-48 (2018). Coistorfor Louron on A. A. et al. A molecular basis for familial hypertrankia | | | | |
| 598 500 | 4. | cardiomyonathy: a beta cardiac myosin heavy chain gene missense mutation. <i>Call</i> 62 | | | | |
| 599 600 | | 999-1006 (1990) | | | | |
| 601 | 5. | Tyska, M.J., <i>et al.</i> Single-molecule mechanics of R403O cardiac myosin isolated from | | | | |
| 602 | | the mouse model of familial hypertrophic cardiomyopathy. <i>Circ Res</i> 86 , 737-744 (2000). | | | | |
| 603 | 6. | Sarkar, S.S., et al. The hypertrophic cardiomyopathy mutations R403Q and R663H | | | | |
| 604 | | increase the number of myosin heads available to interact with actin. Sci Adv 6, eaax0069 | | | | |
| 605 | _ | (2020). | | | | |
| 606 607 | 7. | Gaudelli, N.M., <i>et al.</i> Programmable base editing of A*T to G*C in genomic DNA without DNA cleavage. <i>Nature</i> 551 , 464-471 (2017). | | | | |
| | | | | | | |

| 608 | 8. | Komor, A.C., Kim, Y.B., Packer, M.S., Zuris, J.A. & Liu, D.R. Programmable editing of |
|-----|-----|------------------------------------------------------------------------------------------------|
| 609 | | a target base in genomic DNA without double-stranded DNA cleavage. Nature 533, 420- |
| 610 | | 424 (2016). |
| 611 | 9. | Koblan, L.W., et al. In vivo base editing rescues Hutchinson-Gilford progeria syndrome |
| 612 | | in mice. <i>Nature</i> 589 , 608-614 (2021). |
| 613 | 10. | Suh, S., et al. Restoration of visual function in adult mice with an inherited retinal disease |
| 614 | | via adenine base editing. Nat Biomed Eng 5, 169-178 (2021). |
| 615 | 11. | Chemello, F., et al. Precise correction of Duchenne muscular dystrophy exon deletion |
| 616 | | mutations by base and prime editing. Sci Adv 7(2021). |
| 617 | 12. | Reichart, D., et al. Efficient in vivo Genome Editing Prevents Hypertrophic |
| 618 | | Cardiomyopathy in Mice. Nature Medicine (2022). |
| 619 | 13. | Koblan, L.W., et al. Improving cytidine and adenine base editors by expression |
| 620 | | optimization and ancestral reconstruction. Nat Biotechnol 36, 843-846 (2018). |
| 621 | 14. | Richter, M.F., et al. Phage-assisted evolution of an adenine base editor with improved |
| 622 | | Cas domain compatibility and activity. Nat Biotechnol 38, 883-891 (2020). |
| 623 | 15. | Walton, R.T., Christie, K.A., Whittaker, M.N. & Kleinstiver, B.P. Unconstrained genome |
| 624 | | targeting with near-PAMless engineered CRISPR-Cas9 variants. Science 368, 290-296 |
| 625 | | (2020). |
| 626 | 16. | Nishimasu, H., et al. Engineered CRISPR-Cas9 nuclease with expanded targeting space. |
| 627 | | <i>Science</i> 361 , 1259-1262 (2018). |
| 628 | 17. | Kleinstiver, B.P., et al. High-fidelity CRISPR-Cas9 nucleases with no detectable |
| 629 | | genome-wide off-target effects. Nature 529, 490-495 (2016). |
| 630 | 18. | Marian, A.J. & Braunwald, E. Hypertrophic Cardiomyopathy: Genetics, Pathogenesis, |
| 631 | | Clinical Manifestations, Diagnosis, and Therapy. Circ Res 121, 749-770 (2017). |
| 632 | 19. | Concordet, J.P. & Haeussler, M. CRISPOR: intuitive guide selection for CRISPR/Cas9 |
| 633 | | genome editing experiments and screens. Nucleic Acids Res 46, W242-W245 (2018). |
| 634 | 20. | Pua, C.J., et al. Genetic Studies of Hypertrophic Cardiomyopathy in Singaporeans |
| 635 | | Identify Variants in TNNI3 and TNNT2 That Are Common in Chinese Patients. Circ |
| 636 | | Genom Precis Med 13, 424-434 (2020). |
| 637 | 21. | Toepfer, C.N., et al. Myosin Sequestration Regulates Sarcomere Function, |
| 638 | | Cardiomyocyte Energetics, and Metabolism, Informing the Pathogenesis of Hypertrophic |
| 639 | | Cardiomyopathy. Circulation 141, 828-842 (2020). |
| 640 | 22. | Cohn, R., et al. A Contraction Stress Model of Hypertrophic Cardiomyopathy due to |
| 641 | | Sarcomere Mutations. Stem Cell Reports 12, 71-83 (2019). |
| 642 | 23. | Vakrou, S. & Abraham, M.R. Hypertrophic cardiomyopathy: a heart in need of an energy |
| 643 | | bar? Front Physiol 5, 309 (2014). |
| 644 | 24. | Lyons, G.E., Schiaffino, S., Sassoon, D., Barton, P. & Buckingham, M. Developmental |
| 645 | | regulation of myosin gene expression in mouse cardiac muscle. J Cell Biol 111, 2427- |
| 646 | | 2436 (1990). |
| 647 | 25. | Geisterfer-Lowrance, A.A., et al. A mouse model of familial hypertrophic |
| 648 | | cardiomyopathy. Science 272, 731-734 (1996). |
| 649 | 26. | Ma, S., et al. Efficient Correction of a Hypertrophic Cardiomyopathy Mutation by |
| 650 | | ABEmax-NG. Circ Res 129, 895-908 (2021). |
| 651 | 27. | Ishikawa, K., Weber, T. & Hajjar, R.J. Human Cardiac Gene Therapy. Circ Res 123, 601- |
| 652 | | 613 (2018). |
| | | |

| 653 | 28. | Zettler, J., Schutz, V. & Mootz, H.D. The naturally split Npu DnaE intein exhibits an |
|-----|-----|-----------------------------------------------------------------------------------------------------|
| 654 | | extraordinarily high rate in the protein trans-splicing reaction. FEBS Lett 583, 909-914 |
| 655 | | (2009). |
| 656 | 29. | Prasad, K.M., Xu, Y., Yang, Z., Acton, S.T. & French, B.A. Robust cardiomyocyte- |
| 657 | | specific gene expression following systemic injection of AAV: in vivo gene delivery |
| 658 | | follows a Poisson distribution. <i>Gene Ther</i> 18 , 43-52 (2011). |
| 659 | 30. | Cellular, T., and Gene Therapies Advisory Committee (CTGTAC) Meeting #70. Toxicity |
| 660 | 201 | Risks of Adeno-associated Virus (AAV) Vectors for Gene Therapy. (ed. (FDA). |
| 661 | | F.a.D.A.) (FDA-2021-N-0651, 2021). |
| 662 | 31. | Teekakirikul, P., et al. Cardiac fibrosis in mice with hypertrophic cardiomyopathy is |
| 663 | | mediated by non-myocyte proliferation and requires Tgf-beta. J Clin Invest 120 , 3520- |
| 664 | | 3529 (2010). |
| 665 | 32. | Pinto, A.R., et al. Revisiting Cardiac Cellular Composition. Circ Res 118, 400-409 |
| 666 | | (2016). |
| 667 | 33. | Jiang, J., Wakimoto, H., Seidman, J.G. & Seidman, C.E. Allele-specific silencing of |
| 668 | | mutant Myh6 transcripts in mice suppresses hypertrophic cardiomyopathy. <i>Science</i> 342 , |
| 669 | | 111-114 (2013). |
| 670 | 34. | Pare, J.A., Fraser, R.G., Pirozynski, W.J., Shanks, J.A. & Stubington, D. Hereditary |
| 671 | | cardiovascular dysplasia. A form of familial cardiomyopathy. Am J Med 31, 37-62 |
| 672 | | (1961). |
| 673 | 35. | Fananapazir, L. & Epstein, N.D. Genotype-phenotype correlations in hypertrophic |
| 674 | | cardiomyopathy. Insights provided by comparisons of kindreds with distinct and identical |
| 675 | | beta-myosin heavy chain gene mutations. <i>Circulation</i> 89 , 22-32 (1994). |
| 676 | 36. | Ho, C.Y., et al. Genotype and Lifetime Burden of Disease in Hypertrophic |
| 677 | | Cardiomyopathy: Insights from the Sarcomeric Human Cardiomyopathy Registry |
| 678 | | (SHaRe). Circulation 138, 1387-1398 (2018). |
| 679 | 37. | Carroll, K.J., et al. A mouse model for adult cardiac-specific gene deletion with |
| 680 | | CRISPR/Cas9. Proc Natl Acad Sci US A 113, 338-343 (2016). |
| 681 | 38. | Maron, B.J., Yeates, L. & Semsarian, C. Clinical challenges of genotype positive (+)- |
| 682 | | phenotype negative (-) family members in hypertrophic cardiomyopathy. Am J Cardiol |
| 683 | | 107 , 604-608 (2011). |
| 684 | 39. | Green, E.M., et al. A small-molecule inhibitor of sarcomere contractility suppresses |
| 685 | | hypertrophic cardiomyopathy in mice. Science 351, 617-621 (2016). |
| 686 | 40. | Stern, J.A., et al. A Small Molecule Inhibitor of Sarcomere Contractility Acutely |
| 687 | | Relieves Left Ventricular Outflow Tract Obstruction in Feline Hypertrophic |
| 688 | | Cardiomyopathy. <i>PLoS One</i> 11 , e0168407 (2016). |
| 689 | 41. | Ladage, D., Ishikawa, K., Tilemann, L., Muller-Ehmsen, J. & Kawase, Y. Percutaneous |
| 690 | | methods of vector delivery in preclinical models. Gene Ther 19, 637-641 (2012). |
| 691 | 42. | Cheng, Q., et al. Selective organ targeting (SORT) nanoparticles for tissue-specific |
| 692 | | mRNA delivery and CRISPR-Cas gene editing. Nat Nanotechnol 15, 313-320 (2020). |
| 693 | 43. | Banskota, S., et al. Engineered virus-like particles for efficient in vivo delivery of |
| 694 | | therapeutic proteins. Cell 185, 250-265 e216 (2022). |
| 695 | 44. | Tabebordbar, M., et al. Directed evolution of a family of AAV capsid variants enabling |
| 696 | | potent muscle-directed gene delivery across species. Cell 184, 4919-4938 e4922 (2021). |
| 697 | 45. | Weinmann, J., et al. Identification of a myotropic AAV by massively parallel in vivo |
| 698 | | evaluation of barcoded capsid variants. Nat Commun 11, 5432 (2020). |

| 699 | 46. | Lompre, A.M., et al. Species- and age-dependent changes in the relative amounts of |
|-----|------|------------------------------------------------------------------------------------------------|
| 700 | | cardiac myosin isoenzymes in mammals. Dev Biol 84, 286-290 (1981). |
| 701 | 47. | Desai, M.Y., et al. Study design and rationale of VALOR-HCM: evaluation of |
| 702 | | mavacamten in adults with symptomatic obstructive hypertrophic cardiomyopathy who |
| 703 | | are eligible for septal reduction therapy. Am Heart J 239, 80-89 (2021). |
| 704 | 48. | Saberi, S., et al. Mavacamten Favorably Impacts Cardiac Structure in Obstructive |
| 705 | | Hypertrophic Cardiomyopathy: EXPLORER-HCM Cardiac Magnetic Resonance |
| 706 | | Substudy Analysis. Circulation 143, 606-608 (2021). |
| 707 | 49. | FDA, U. CAMZYOS (mavacamten) capsules for oral use. (2022). |
| 708 | 50. | Ho, C.Y., et al. Evaluation of Mavacamten in Symptomatic Patients With Nonobstructive |
| 709 | | Hypertrophic Cardiomyopathy. J Am Coll Cardiol 75, 2649-2660 (2020). |
| 710 | 51. | Ran, F.A., et al. Genome engineering using the CRISPR-Cas9 system. Nat Protoc 8, |
| 711 | | 2281-2308 (2013). |
| 712 | 52. | Huang, T.P., et al. Circularly permuted and PAM-modified Cas9 variants broaden the |
| 713 | | targeting scope of base editors. Nat Biotechnol 37, 626-631 (2019). |
| 714 | 53. | Levy, J.M., et al. Cytosine and adenine base editing of the brain, liver, retina, heart and |
| 715 | | skeletal muscle of mice via adeno-associated viruses. Nat Biomed Eng 4, 97-110 (2020). |
| 716 | 54. | Burridge, P.W., et al. Chemically defined generation of human cardiomyocytes. Nat |
| 717 | | <i>Methods</i> 11 , 855-860 (2014). |
| 718 | 55. | Correia, C., et al. Distinct carbon sources affect structural and functional maturation of |
| 719 | | cardiomyocytes derived from human pluripotent stem cells. Sci Rep 7, 8590 (2017). |
| 720 | 56. | Kluesner, M.G., et al. EditR: A Method to Quantify Base Editing from Sanger |
| 721 | | Sequencing. CRISPR J 1, 239-250 (2018). |
| 722 | 57. | Atmanli, A., et al. Cardiac Myoediting Attenuates Cardiac Abnormalities in Human and |
| 723 | | Mouse Models of Duchenne Muscular Dystrophy. Circ Res 129, 602-616 (2021). |
| 724 | 58. | Kijlstra, J.D., et al. Integrated Analysis of Contractile Kinetics, Force Generation, and |
| 725 | | Electrical Activity in Single Human Stem Cell-Derived Cardiomyocytes. Stem Cell |
| 726 | | <i>Reports</i> 5, 1226-1238 (2015). |
| 727 | 59. | Doench, J.G., et al. Optimized sgRNA design to maximize activity and minimize off- |
| 728 | | target effects of CRISPR-Cas9. Nat Biotechnol 34, 184-191 (2016). |
| 729 | 60. | Clement, K., et al. CRISPResso2 provides accurate and rapid genome editing sequence |
| 730 | | analysis. Nat Biotechnol 37, 224-226 (2019). |
| 731 | 61. | Miura, H., Quadros, R.M., Gurumurthy, C.B. & Ohtsuka, M. Easi-CRISPR for creating |
| 732 | | knock-in and conditional knockout mouse models using long ssDNA donors. Nat Protoc |
| 733 | | 13 , 195-215 (2018). |
| 734 | 62. | Creed, H.A. & Tong, C.W. Preparation and Identification of Cardiac Myofibrils from |
| 735 | | Whole Heart Samples. Methods Mol Biol 2319, 15-24 (2021). |
| 736 | 63. | Cui, M. & Olson, E.N. Protocol for Single-Nucleus Transcriptomics of Diploid and |
| 737 | | Tetraploid Cardiomyocytes in Murine Hearts. STAR Protoc 1, 100049 (2020). |
| 738 | | |
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755 Author contributions:

A.C.C., R.B.D., and E.N.O. conceived of the project and designed the experiments. A.C.C., F.C.,

H.L., and A.A. conducted in vitro iPSC-CM experiments. A.C.C., M.C., F.C., H.L., and Y.Z.

conducted in vivo experiments. W.T. performed mouse echocardiography. J.M. performed

mouse zygote injections. K.C., A.C.C., and L.X. performed bioinformatics analysis. A.C.C.,

760 N.L., R.B.D., and E.N.O. wrote the manuscript.

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762 **Competing interests:** E.N.O. is a consultant for Vertex Pharmaceuticals and Tenaya

763 Therapeutics. The other authors declare that they have no competing interests.



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Fig. 1: In vitro optimization of ABE system to correct a pathogenic MYH7 variant. a, A 765 candidate sgRNA, h403 sgRNA, for base editing to correct the MYH7 c.1208G>A (p.R403Q) 766 missense variant. Base editing could convert the neutrally charged glutamine pathogenic variant 767 back to a positively charged arginine, restoring proper function of the myosin head. b, Eight 768 candidate base editor variants were screened for their efficiencies in correcting the pathogenic 769 adenine to a guanine using the candidate h403 sgRNA within a homozygous MYH7 c.1208G>A 770 iPSC line (HD^{403/403}). c, DNA editing efficiency of all adenines within the target protospacer in 771 $HD^{403/403}$ iPSCs 72 h post-transfection with candidate base editors. Data are mean \pm s.d. across 772 three technical replicates. Numbering is with the first base 5' of the PAM as 1; target pathogenic 773 adenine is position A16. 774



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Fig. 2: Analyses of HCM patient-derived iPSC-CM function upon base editing correction. a, 776 Workflow for reprogramming iPSCs from a healthy donor (HD) and two HCM patients (HCM1 777 and HCM2), followed by pathogenic variant knock-in for the HD line, and base editing correction 778 for the HCM1 and HCM2 lines. Isogenic clonal lines were then isolated and differentiated into 779 CMs for downstream analyses of iPSC-CM function. b, Deep-High throughput sequencing (HTS) 780 to measure editing for all adenines within the on-target protospacer. Target pathogenic adenine is 781 A16. Deep sequencingHTS was performed for ABE-treated MYH7^{403/+}-HCM1^{403/+}, and MYH7^{403/+} 782 HCM2^{403/+} iPSCs. c, Quantification of peak systolic force of MYH7^{403/+} and MYH7^{WT} iPSC-CMs 783 from HD, HCM1, and HCM2 patients for indicated cell lines. (n=31-57 for each genotype) Data 784 are mean \pm s.e.m. across three separate differentiations. **d**, Oxygen consumption rate (OCR) as a 785

function of time in indicated cell lines following exposure to the electron transport chain complex inhibitors, oligomycin, carbonyl cyanide *m*-chlorophenyl hydrazone (CCCP), and Antimycin A (AntA) (top), and mean and distribution of values across four timepoints for basal OCR (bottom left) and maximal OCR (bottom right) for indicated cell lines. Data are mean \pm s.d. across three separate differentiations. Each data point is from 14-16 Seahorse assay wells. ***P* < 0.01, *****P* < 0.001, *****P* < 0.0001 by Student's unpaired two-sided t-test.





Fig. 3: Generation of a humanized HCM mouse model. a, A humanized HCM mouse model generated was by replacing part of the native murine *Myh6* genomic sequence with the human *MYH7* sequence containing the p.R403Q variant via CRISPR-Cas9 homology-directed repair (HDR). Sanger sequencing chromatograms show the native $Myh6^{WT}$ sequence (top), the humanized $Myh6^{h403/+}$ mouse model sequence (middle), and a human HCM patient-derived iPSC line sequence (bottom). Yellow squares indicate knocked-in SNPs; Sanger sequencing shows

double nucleotide peaks for SNPs. b, Gross histology (top), and Masson's trichrome staining of
coronal (4-chamber) (middle) and transverse (bottom) sections of the humanized mouse model for
the wildtype (left), heterozygous (middle), and homozygous (right) genotypes at postnatal day 8.
Scale bar, 1 mm. c, Masson's trichrome, Picrosirius red, and hematoxylin & eosin staining of heart
sections of the humanized mouse model for the wildtype (left) and heterozygous (right) genotypes
at 9 months of age. Scale bar, 1 mm for 10x images top, 100 µm for 10x images middle, 25 µm
for 40x images bottom.



Fig. 4: Prevention of HCM by dual AAV9 ABE editing of $Myh6^{h403/+}$ mice. a, Schematic of dual AAV9 ABE system encoding ABEmax-VRQR base editor halves and h403_sgRNA to target the human *MYH7* p.R403Q variant. b, Experimental outline for intrathoracic injection of $Myh6^{WTh403/+}$ or $Myh6^{h403/+}$ mice with saline or dual AAV9 ABE at P0 followed by serial

echocardiograms. Chow diet supplemented with 0.1% Cyclosporine A was given at 5 weeks of 812 age for 11 weeks. c-h, Left ventricular anterior wall thickness at diastole (c), left ventricular 813 posterior wall thickness at diastole (d), left ventricular internal diameter at diastole (e) and systole 814 (f), ejection fraction (g), and fractional shortening (h), of $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, or ABE-815 treated $Myh6^{h403/+}$ mice at 16 weeks of age. n=5 for each group. Data are mean \pm s.e.m. i, 816 Representative Masson's trichrome staining of serial (500 µm interval) transverse sections for 817 $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, or ABE-treated $Myh6^{h403/+}$ mice at 16 weeks of age. Scale bar, 1 818 819 mm. j, Ventricular cross-sectional area measurements from n=3-5 mice for each experimental 820 group. Data are mean \pm s.d. **k**, Average wall thickness measurements from n=3-5 mice for each experimental group. Data are mean \pm s.d. **I**, Heart weight (HW) to tibia length (TL) measurements 821 from n=5 mice for each experimental group. Data are mean \pm s.d. **m**, Percentage of collagen area 822 *n*=3-5 mice for each experimental group. Data are mean \pm s.d. **P* < 0.05, ***P* < 0.01 by Student's 823 unpaired two-sided *t*-test. 824





Fig. 5: Genomic and transcriptomic changes following dual AAV9 ABE injection in mice. a, 826 Cardiomyocyte nuclei were isolated from ventricles of 18 weeks old $Myh6^{WT}$ mice, $Myh6^{h403/+}$ 827 mice, or ABE-treated $Myh6^{h403/+}$ mice to assess genomic correction and transcriptomic changes. 828 **b**, DNA-editing efficiency for correcting the pathogenic adenine nucleotide following dual AAV9 829 ABE treatment. Data are mean \pm s.d. c, Percentage of expressed pathogenic transcripts in ABE-830 treated $Myh6^{h403/+}$ mice compared to $Myh6^{h403/+}$ mice. Data are mean \pm s.d. **d**, Bystander editing 831 in ABE-treated $Myh6^{h403/+}$ mice compared to saline-treated mice. Data are mean \pm s.d. e, 832 Transcriptome-wide nuclear levels of A-to-I RNA editing in $Mvh6^{WT}$ mice, $Mvh6^{h403/+}$ mice, and 833 ABE-treated $Myh6^{h403/+}$ mice. Data are mean \pm s.d. **f**, Heat map of 257 differentially expressed 834 genes amongst $Myh6^{WT}$ or $Myh6^{h403/+}$ mice and ABE-treated $Myh6^{h403/+}$ mice. Samples and genes 835 are ordered by hierarchical clustering. Data was scaled by the sum of each row and are displayed 836 as row min and row max. ABE-treated $Myh6^{h403/+}$ mice cluster with $Myh6^{WT}$ mice. Editing 837 efficiency for each ABE-treated *Myh6^{h403/+}* mouse is indicated. **g**, Top gene ontology (GO) terms 838

- associated with the differentially expressed genes in the comparison of h403/+ vs WT (top) and
- 840 h403/+ ABE-treated vs h403/+ (bottom). h, Fold change expression of *Nppa* mRNA expression
- for $Myh6^{h403/+}$ mice and ABE-treated $Myh6^{h403/+}$ mice normalized to $Myh6^{WT}$ mice. Data from
- 842 RNA-seq and qPCR. Data are mean \pm s.d. For all: **P* < 0.05 by Student's unpaired two-sided *t*-
- test, n=3 biological replicates for each group.

| | | | Echocardiography | Measurements | | | | | |
|-----|----------------|----------------|-----------------------------------------|--------------|------------------|----------------------|--|--|--|
| | | LVAW;d (mm) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 0.796 ± 0.0453 | 0.908 ± 0.0283 | 0.775 ± 0.0510 | 0.069 | 0.757 | 0.051 | | | |
| 12w | 0.908 ± 0.0434 | 1.07 ± 0.0428 | 0.829 ± 0.0243 | 0.023 | 0.154 0.001 | | | | |
| 16w | 0.883 ± 0.0441 | 1.07 ± 0.0443 | .07 ± 0.0443 0.950 ± 0.0414 0.017 0.299 | | | | | | |
| | | LVID;d (mm) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 2.83 ± 0.126 | 2.54 ± 0.110 | 2.64 ± 0.0749 | 0.069 | 0.757 | 0.051 | | | |
| 12w | 2.90 ± 0.0983 | 2.45 ± 0.0986 | 2.84 ± 0.149 | 0.013 | 0.763 | 0.060 | | | |
| 16w | 2.81 ± 0.0540 | 2.34 ± 0.142 | 2.78 ± 0.110 | 0.015 | 0.038 | | | | |
| | | LVID;s (mm) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 1.27 ± 0.109 | 1.05 ± 0.0580 | 1.20 ± 0.0988 | 0.107 | 0.638 | 0.222 | | | |
| 12w | 1.37 ± 0.0729 | 1.00 ± 0.0544 | 1.16 ± 0.0794 | 0.004 | 0.085 | 0.147 | | | |
| 16w | 1.24 ± 0.0520 | 0.940 ± 0.0713 | 1.21 ± 0.108 | 0.010 | 0.071 | | | | |
| | | LVPW;d (mm) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 0.850 ± 0.0349 | 0.899 ± 0.0262 | 0.771 ± 0.0695 | 0.300 | 0.335 | 0.123 | | | |
| 12w | 0.910 ± 0.0471 | 1.00 ± 0.0605 | 0.807 ± 0.0546 | 0.264 | 0.192 | 0.044 | | | |
| 16w | 0.867 ± 0.0590 | 1.04 ± 0.0809 | 0.970 ± 0.0709 | 0.128 | 0.299 | 0.548 | | | |
| | | EF (%) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 87.0 ± 1.79 | 88.6 ± 0.606 | 86.8 ± 1.86 | 0.428 | 0.922 | 0.374 | | | |
| 12w | 85.8 ± 2.01 | 89.4 ± 1.40 | 88.2 ± 2.13 | 0.188 | 0.448 | 0.652 | | | |
| 16w | 87.4 ± 1.63 | 91.7 ± 0.807 | 87.3 ± 1.61 | 0.045 | 0.965 | 0.039 | | | |
| | | FS (%) | | | P Value | | | | |
| | WT | h403/+ | ABE h403/+ | h403/+ vs WT | ABE h403/+ vs WT | ABE h403/+ vs h403/+ | | | |
| 8w | 55.4 ± 2.19 | 56.7 ± 0.894 | 52.6 ± 3.30 | 0.592 | 0.509 | 0.269 | | | |
| 12w | 53.9 ± 2.40 | 58.1 ± 2.25 | 57.1 ± 2.79 | 0.238 | 0.404 | 0.798 | | | |
| 16w | 55.8 ± 2.33 | 61.6 ± 1.40 | 55.5 ± 2.12 | 0.066 | 0.928 | 0.043 | | | |

Table 1: Summary of echocardiographic measurements for humanized mice. Echocardiographic measurements in $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, and ABE-treated $Myh6^{h403/+}$ mice (8-16 weeks, n=5) for changes in left ventricular anterior wall thickness (LVAW), left ventricular internal diameter at diastole (LVID;d) and at systole (LVID;s), left ventricular posterior wall thickness (LVPW), ejection fraction (EF) and fractional shortening (FS). *P* values are calculated by Student's unpaired two-sided *t*-test for given comparisons. *P* values <0.05 are in bold. Data are mean \pm s.e.m.



Extended Data Fig. 1: Generation of isogenic HD^{403/+} and HD^{403/403} iPSCs by homology-854 directed repair. a, Using iPSCs derived from a healthy donor (HDWT), the MYH7 p.R403Q 855 (c.1208G>A) variant was introduced by CRISPR-Cas9-based homology-directed repair (HDR) 856 using SpCas9, a sgRNA (spacer sequence colored in green, PAM sequence colored in gold), and 857 a single-stranded oligodeoxynucleotide (ssODN) donor template containing the pathogenic 858 variant. A heterozygous genotype (HD^{403/+}) and homozygous genotype (HD^{403/403}) were isolated. 859 860 Chromatograms highlighting insertion of the pathogenic variant and corresponding amino acid changes are shown for indicated genotypes. Red arrows indicate coding nucleotide 1208 and amino 861 acid 403. b, Sanger sequencing chromatogram showing no insertion of the pathogenic variant on 862 the highly homologous MYH6 gene. Red arrow indicates coding nucleotide 1211 and amino acid 863

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- 404. **c**, HD^{WT} and HD^{403/+} iPSCs readily differentiate into CMs. Cardiac troponin I (cTnI, green)
- ⁸⁶⁵ highlights CMs; nuclei (blue) are marked by DAPI (4',6-diamidino-2-phenylindole). Scale bar, 25
- 866 μ m. **d**, Ratio of *MYH7* to *MYH6* gene expression in HD^{WT} and HD^{403/+} iPSC-CMs as measured by
- 867 quantitative PCR. Data are mean \pm s.d.

| Target | gRNA Sequence | | Gene | 20 |) | | | | | | | 10 | | | | | | 1 | Ρ | A |
|-----------|----------------------|-----|----------------------------------|-----|---|---|-------|---|-----|---|---|----|----------|----|---|-----|---|-----|---|---|
| On Target | CCTCAGGTGAAAGTGGGCAA | TGA | MYH7 | C | C | Т | C A | G | G 1 | G | Α | A | ٩G | Т | G | G G | С | A A | Ν | G |
| OT 1 | CCTCGGGTGAAAGTGGGCAA | CGA | MYH6 | 1 | Ι | Ι | 0 | i | 11 | I | Ι | Ι | | Ι | L | | Ι | 11 | С | Ι |
| OT 2 | CCTAAAGAGAAAATGGGCAA | AGA | Intron; CEP57 |] ı | Ι | | A | Α | I | 1 | Ι | L | Α | 1 | L | | Ι | 11 | Α | 1 |
| OT 3 | TCTCAGATGAAAGTGAGCTA | AGA | FRYL | Т | T | T | - | Τ | A | 1 | Ι | L | | 1 | 1 | 4 | Т | ΤI | Α | 1 |
| OT 4 | CATCAAGTGAAAGTGGACAG | GGA | Intron; SMPDL3B/RP11-460I13.2 | 1 | А | T | | Α | | I | Ι | L | | Ι | I | A | 1 | G | G | Τ |
| OT 5 | CCTCAGGAGAAGATGGACAA | AGA | Intergenic; RP11-27814.2-COLEC10 | 1 | Ι | T | | Ι | I A | 1 | Ι | 1 | G A | I. | Ι | A | 1 | ΙI | Т | Τ |
| OT 6 | TATCAGGTGAAGGTAGGCAA | TGA | STAU2 | Т | Α | Т | | Ι | | I | Ι | 1 | G | Т | Α | | Ι | 11 | Т | Т |
| OT 7 | GCTCAGGAGAAGGTGGACAA | TGA | RP6-127F18.2 | G | 1 | L | | Ι | 4 | 1 | Τ | 1 | G | Ι | L | A | 1 | I I | Т | Ι |
| OT 8 | TCTCAAGGGAGAGTGGGCAA | GGA | Intron; FERMT1-TARDBPP1 | Т | 1 | Ι | | А | 1 0 | i | 1 | G | | Ι | L | | I | 11 | G | Τ |



Extended Data Fig. 2: Computationally determined off-target sites for h403 sgRNA with 869 ABEmax-VRQR. a, Genomic loci of eight candidate off-target sequences (left) and alignments 870 of eight candidate off-target sites to the on-target protospacer (right). Nucleotides that match the 871 protospacer are indicated with a vertical dash. Nucleotides that differ are shown for each site. 872 Numbering of nucleotides in protospacer starts with the nucleotide immediately 5' of the PAM as 873 nucleotide 1. **b**, Deep-High throughput sequencing (HTS) to measure editing for all 58 adenines 874 within the protospacers of the top eight CRISPOR-identified candidate off-target loci. Deep 875 sequencing-HTS was performed for ABE-treated $MYH7^{403/+}$ HCM1, and $MYH7^{403/+}$ HCM2 iPSCs. 876 877

| Mouse Human | DADKSAYLMGLNSADLLKGLCHPQVKVGNEYVTKGQS 37 EADKSAYLMGLNSADLLKGLCHPQVKVGNEYVTKGQN 37 :************************************ | |
|----------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------|
| Mouse Human | $\begin{array}{c ccccccccccccccccccccccccccccccccccc$ | 60 60 |
| Mouse Human | C H P Q V K V G N E Y V T K G Q S TGTCACCCTCAGGTGAAGGTGGGGAACGAGTATGTCACCAAGGGGCAGAGT 111 TGCCACCCTCAGGTGAAAGTGGGCAATGAGTACGTCACCAAGGGGCAGAAT 111 C H P Q V K V G N E Y V T K G Q N *** *** **************************** | |

879 Extended Data Fig. 3: Comparison of mouse and human myosin heavy chain sequences.

880 Homology comparison for mouse α -myosin heavy chain (*Myh6*) and human β -myosin heavy chain

(MYH7) at the amino acid level (top) and DNA sequence level (bottom) around glutamine 403.

882 The h403_sgRNA is illustrated in green and the PAM sequence is illustrated in yellow. The

pathogenic c.1208 G>A variant is located at position 16 within the canonical base editing window

of positions 14-17, counting the adenine nucleotide immediately 5' of the PAM as position 1.



Extended Data Fig. 4: Validation of a dual AAV9 ABE system in mice. a, Injection details for 887 treating $Myh6^{h403/h403}$ mice with ABE-AAV9 or saline. **b**, Kaplan-Meier curve for $Myh6^{WT}$ mice 888 (n=7), $Myh6^{h403/+}$ mice (n=8), $Myh6^{h403/h403}$ mice (n=6), and ABE-treated $Myh6^{h403/h403}$ mice at a 889 low (AAV LOW, n=3) or high dose (AAV HIGH, n=5). Median lifespans: $Myh6^{WT}$ and $Myh6^{h403/+}$ 890 mice, >40 days; $Myh6^{h403/h403}$ mice, 7 days; AAV LOW $Myh6^{h403/h403}$ mice, 9 days (1.3-fold longer, 891 P < 0.05; AAV HIGH *Myh6*^{h403/h403} mice, 15 days (2.1-fold longer, P < 0.01). c, Sanger 892 sequencing chromatograms for a $Myh6^{h403/h403}$ mouse and a AAV HIGH $Myh6^{h403/h403}$ mouse 893 showing 35% on-target editing of the target pathogenic adenine at the cDNA level. d, Four-894

chamber sectioning and Masson's trichrome staining of a AAV HIGH $Myh6^{h403/h403}$ mouse at 15

896 days old of age. *P < 0.05, **P < 0.01 by Mantel-Cox test.



Extended Data Fig. 5: Serial echocardiograms following dual AAV9 ABE editing of 899 $Myh6^{h403/+}$ mice. a-f, Left ventricular anterior wall thickness at diastole (a) left ventricular 900 posterior wall thickness at diastole (b), left ventricular internal diameter at diastole (c) and systole 901 (d), ejection fraction (e), and fractional shortening (f), of $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, or ABE-902 treated $Myh6^{h403/+}$ mice from 8-16 weeks of age. n=5 for each group. Data are mean \pm s.e.m. *P <903 0.05, **P < 0.01 by Student's unpaired two-sided *t*-test for $Myh6^{WT}$ mice compared to $Myh6^{h403/+}$ 904 mice (black) and ABE-treated Myh6^{h403/+} mice compared to Myh6^{h403/+} mice (green). g, 905 Representative M-mode images for $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, and ABE-treated $Myh6^{h403/+}$ 906 mice at 16 weeks of age. 907



Extended Data Fig. 6: Genomic and proteomic analysis of select tissues following dual AAV ABE editing. a, Viral copy numbers for the N terminal AAV and C terminal AAV were quantified from the right atrium (RA), right ventricle (RV), left atrium (LA), left ventricle (LV), lung, liver, spleen, and quadriceps muscle (Quad) from ABE-treated $Myh6^{h403/+}$ mice at 16 weeks of age. b, The percentage of A to G editing was determined by high throughput sequencing (HTS) of genomic DNA in the RA, RV, LA, LV, lung, liver, spleen, and Quad from ABE-treated and saline-

injected $Myh6^{h403/+}$ mice. **c**, The percentage decrease in mutant transcripts in the RA, RV, LA, and 915 LV was determined by HTS of cDNA from ABE-treated and saline-injected $Myh6^{h403/+}$ mice. d, 916 Cardiac myofibrils were isolated from $Myh6^{WT}$ mice, $Myh6^{h403/+}$ mice, and ABE-treated $Myh6^{h403/+}$ 917 mice, run on a 4-20% polyacrylamide gel, and stained with Coomassie G-250. Key sarcomeric 918 proteins are marked, including titin, myosin heavy chain (MHC), myosin binding protein C 919 (MyBP-C), actin, cardiac troponin T (cTnT), cardiac tropomyosin (cTm), and cardiac troponin I 920 (cTnI). Sizes for ladder markings are in kDa. Relative protein amounts for each key sarcomeric 921 protein are normalized to WT. Data are mean \pm s.d. **P* < 0.05 by Student's unpaired two-sided *t*-922 test, *n*=3 biological replicates for each group. 923



926 Extended Data Fig. 7: RNA-sequencing analysis of dual AAV9 ABE editing of $Myh6^{h403/+}$ 927 mice. Volcano plot showing fold-change and *p*-value of genes up-regulated (red) and down-

- regulated (blue) in $Myh6^{h403/+}$ mice compared to $Myh6^{WT}$ mice (top), ABE-treated $Myh6^{h403/+}$ mice compared to $Myh6^{h403/+}$ mice (middle), and ABE-treated $Myh6^{h403/+}$ mice compared to $Myh6^{WT}$
- 930 mice (bottom).
- 931



Supplementary Figure 1. Representative flow cytometry gating strategy for GFP+ iPSCs.
For generation of isogenic iPSCs containing the *MYH7* c.1208 G>A (p.R403Q) mutation via
homology-directed repair, and for base editing correction of the *MYH7* c.1208 G>A (p.R403Q)
mutation, iPSCs were nucleofected with a single plasmid encoding for Cas9 nuclease or an
adenine base editor linked to GFP via a 2A self-cleaving peptide, and the sgRNA of interest.
Cells that were successfully nucleofected with the gene editing components express GFP, and
single GFP+ iPSCs were collected.





944 Supplementary Table 1. Summary of oligos

| Oligo Name | Oligo Sequence |
|---------------------------------------------------------------|-----------------------------------------------------------------|
| sgRNA for HDR | TCATTGCCCACTTTCACCCG |
| Knock-In of | |
| MYH7 R403Q | |
| ssODN for HDR | |
| MVH7 R4030 | GAGTACGTCACCAAGGGGCAG |
| Sequencing for | ACCTCCACATCCTGGGTTCAA |
| hMYH7 F | |
| Sequencing for | GTGGAGGAGAGACCCATATT |
| hMYH/R Sequencing for | |
| hMYH6 F | UCAUCE IN FAUTUAUCCAAU |
| Sequencing for | AGGAGCAAGCGAGTGATTGT |
| h403 sgRNA | CCGCAGGTGAAAGTGGGCAA |
| HTS ON-Target | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGTCCTCTCATACACTGCCTTGG |
| F | |
| HTS ON-Target | GTCTCGTGGGGCTCGGAGATGTGTATAAGAGACAGCACCATGCCTGGCTAATTTT |
| HTS OFF1 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGGGACAATGACTGCCTCTGT |
| HTS OFF1 R | GTCTCGTGGGGCTCGGAGATGTGTATAAGAGACAGTACCTCATGGGGCTGAACTC |
| HTS OFF2 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCAGGTCTCGATTCCAAGGAG |
| HTS OFF2 R | GTCTCGTGGGGCTCGGAGATGTGTATAAGAGACAGGCACAACCCACAAGTTTGTT T |
| HTS OFF3 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGTTTTCAAAATATTCCTGCTCAC T |
| HTS OFF3 R | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGAGGCACCTTTCTGTGTGCTT |
| HTS OFF4 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGATTCTGGATGCAGGATTTGC |
| HTS OFF4 R | GTCTCGTGGGGCTCGGAGATGTGTATAAGAGACAGGTGGACAACAGGCCACTCTT |
| HTS OFF5 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGGACAATTTGTATTTAGCTTA TTTTC |
| HTS OFF5 R | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGTCCCCTGCTTTTCTCTGTGT |
| HTS OFF6 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGTGATCCTGAAGATTAGTGGAT GC |
| HTS OFF6 R | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCCATCCTGAGATAATCCTCC |
| HTS OFF7 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGACCTAGGAGGCTGGGATTGT |
| HTS OFF7 R | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCATGACAAGGAGTCCGAGGT |
| HTS OFF8 F | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGCCCCTGGTTACAGCATAAG |
| HTS OFF8 R | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCCACAACCACTGACTG |
| sgRNA for Knock-In of MYH7 R403Q into murine Myh6 | TCGTTCCCCACCTTCACCCG |

| ssODN for | TGGGACAAAGGAATGGAGGTACTGAAAATGCTTCCCCTCTCCTTGTCTATCAGAT |
|----------------|---------------------------------------------------------|
| Knock-In of | GCTGACAAATCAGCCTACCTCATGGGGGCTGAACTCAGCCGACCTGCTCAAGGGG |
| MYH7 R403Q | CTGTGCCACCCTCAGGTGAAAGTGGGCAATGAGTACGTCACCAAGGGGCAGAGT |
| into murine | GTACAGCAAGTGTACTAT |
| Myh6 | |
| Genotyping for | GAGAAGCAGTGGTCATCATC |
| Myh6 F | |
| Genotyping for | GTGAGAAACACGTGGTGTCC |
| Myh6 R | |
| HTS Myh6 On- | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGGATCAAGGACATGGCAAAT |
| Target F | |
| HTS Myh6 On- | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGCTTGGTCTCCAGGGTTG |
| Target R | |
| HTS Myh6 | TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGATGGCACAGAAGATGCTGA |
| cDNA On-Target | |
| F | |
| HTS Myh6 | GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCGAACATGTGGTGGTTGAAG |
| cDNA On-Target | |
| R | |
| Sanger Myh6 | GCTCTTGGCCACTGATAGTGC |
| cDNA On-Target | |
| F | |
| Sanger Myh6 | GCTCAAAGCTGTTGAAATCG |
| cDNA On-Target | |
| R | |
| VCN N terminal | ACCAGAAAGAGCGAGGAAAC |
| AAV half F | |
| VCN N terminal | TCGTTGGGCAGGTTCTTATC |
| AAV half R | |
| VCN N terminal | /56-FAM/TTGGTCATC/ZEN/CGCTCGATGAAGCTC/3IABkFQ/ |
| AAV Probe | |
| VCN C terminal | CCCAAGAGGAACAGCGATAAG |
| AAV half F | |
| VCN C terminal | CCACCAGCACAGAATAG |
| AAV half R | |
| VCN C terminal | /56-FAM/ATCGCCAGA/ZEN/AAGAAGGACTGGGAC/3IABkFQ/ |
| AAV Probe | |