

Single-cell full-length spatial transcriptomics reveal the unexplored isoform diversity of the myocardium post-MI

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2 ABSTRACT

We introduce Single-cell Nanopore Spatial Transcriptomics (scNAST), a software suite to facilitate the analysis of spatial gene expression from second- and third-generation sequencing, allowing to generate a full-length single-cell transcriptional landscape of the tissue microenvironment. Taking advantage of the Visium Spatial platform, we adapted a strategy recently developed to assign barcodes to long-read single-cell sequencing data for spatial capture technology. Here, we demonstrate our workflow using four short axis sections of the mouse heart following myocardial infarction. We constructed a *de novo* transcriptome using long-read data, and successfully assigned 19,794 transcript isoforms in total, including clinically-relevant, but yet uncharacterized modes of transcription, such as intron retention or antisense overlapping transcription. We showed a higher transcriptome complexity in the healthy regions, and identified intron retention as a mode of transcription associated with the infarct area. Our data revealed a clear regional isoform switching among differentially used transcripts for genes involved in cardiac muscle contraction and tissue morphogenesis. Molecular signatures involved in cardiac remodeling integrated with morphological context may support the development of new therapeutics towards the treatment of heart failure and the reduction of cardiac complications.

18 **Keywords:** Spatial transcriptomics, Single-cell RNA sequencing, Oxford Nanopore Technology, Myocardial Infarction

INTRODUCTION

Cell type heterogeneity has recently emerged as a major aspect in redrawing the cellular picture of the mammalian heart (Wang et al., 2020; Tucker et al., 2020; Litviňuková et al., 2020). Single-cell RNA-seq (scRNA-seq) technology has enabled to explore crosstalk of different cardiac cell populations to identify response signatures involved in remodeling after myocardial infarction (MI) and ischemic injury (Cui et al., 2020; Forte et al., 2020; Ruiz-Villalba et al., 2020; Vafadarnejad et al., 2020; Molenaar et al., 2021; Gladka et al., 2021; Tombor et al., 2021; Heinrichs et al., 2021). These and other data provide a valuable compendium of information to better understand transcriptional changes occurring in cardiomyocyte and non-cardiomyocyte sub-populations in healthy, injured, and regenerating hearts. However, in all these studies, the original tissue architecture is destroyed and, in general, the morphological context is lost, including the relationship of cells to infarct, border and remote zones (van Duijvenboden et al., 2019a).

Recent development in spatial transcriptomics addresses this challenge, but few studies only have provided spatially resolved insights into the cardiac transcriptome. Techniques such as microdissection (Wu et al., 2016; Burkhard and Bakkers, 2018) or, in particular, spatially barcoded arrays and *in situ* capturing, have enabled to retain the spatial information while profiling the whole transcriptome at near single-cell resolution, allowing to shed light on localized tissue neighbourhoods (Asp et al., 2017, 2019), but a detailed characterization of cellular zones of injury, repair and/or remodeling is lacking. MI is a complex spatio-temporal heterogeneous disease involving the whole heart, and unbiased spatial transcriptomics holds a promise to add tissue context to molecular profiling in the search of novel therapeutics.

One of the most established spatial transcriptomics methods is now widely available as the Visium platform by 10x Genomics. After the tissue section is fixed on the spatial slide, stained, and imaged, it is permeabilized to release RNA to bind to capture probes for on-slide cDNA synthesis. Library preparation is performed off-slide, and spatial barcodes and tissue image are used to overlay transcriptomics data with tissue context. Although several new such methods have recently been published (Rodrigues et al., 2019; Liu et al., 2020), they are for the most part relying on short-read library preparation. Thus they fail to generate sufficient overlaps to reconstruct transcriptomes *de novo*.

Adequate gene and transcript models are instrumental towards relevant proof of concepts and investigational new drug development in translational cardiac research (Müller et al., 2021). Third-generation or long-read sequencing allow to reconstruct truthful assemblies with fewer gaps and to characterize complete transcript isoforms and chimeric transcripts. This has recently been taken to the single-cell level using either Pacific Biosciences (PacBio) or Oxford Nanopore technology (Gupta et al., 2018; Lebrigand et al., 2020; Volden and Vollmers, 2021; Wang et al., 2021; Joglekar et al., 2021).

Here, taking advantage of the conceptual similarity between spatial and cell barcodes, we introduce Single-cell Nanopore Spatial Transcriptomics (SCNAST), a set of tools to facilitate whole transcriptome spatial profiling of full-length transcripts, based on a previously published Bayesian approach for cell barcode assignment (Wang et al., 2021). Our method relies on the commercially available Visium platform by 10x Genomics and Nanopore long-read sequencing, although it can in principle be extended to other technologies, as long as a hybrid sequencing approach is used for spatial barcode assignment.

In this short report, we demonstrate how SCNAST can be used to characterize the spatial transcriptional landscape of the heart post-MI. Using our workflow, we were able to assign a total of 7,616 spatial spots across four short axis sections of the heart, corresponding to distinct 19,794 transcript isoforms in total, encoded by 12,474 genes. We showed a higher transcriptome complexity in the healthy regions, and identified intron retention as a mode of transcription associated with the injured areas. Our results showed a

61 clear regional isoform switching among differentially used transcripts for genes involved in cardiac muscle
62 contraction and tissue morphogenesis, many of them clinically relevant, opening new opportunities in
63 translational cardiac research.

MATERIAL AND METHODS

64 Experimental model of myocardial infarction

65 A C57BL/6 mouse (female, 8 weeks old, Janvier Labs) was exposed to 5% isoflurane for anesthesia. An
66 intubation cannula was inserted orally into the trachea. The mouse was fixed on a heating plate at 37°C
67 and maintained under anesthesia with 2% isoflurane. An incision was made from the left sternum to the
68 midclavicular line. Skins and muscle layers were stretched with forceps and sutures. Another incision was
69 made between the third and fourth intercostal space. The heart was exposed and subjected to permanent
70 myocardial infarction with ligation of the left anterior descending (LAD) coronary artery. When the ribs
71 and skins were fully closed, the isoflurane supply was cut off. Oxygen was then supplied until normal
72 breathing was resumed.

73 Heart extraction and cryosection

74 Three days after permanent LAD ligation, the mouse was sacrificed for organ harvest. After washing with
75 cold PBS several times to remove the blood, the heart was transferred into a bath of isopentane (Millipore
76 Sigma) frozen by liquid nitrogen. The freshly obtained heart was kept fully submerged in isopentane
77 for 5 min until fully frozen. Pre-cooled Cryomold on the dry ice was filled with chilled TissueTek OCT
78 compound without introducing bubbles. The frozen tissue was then transferred into the OCT with pre-
79 cooled forceps and placed on the dry ice until the OCT was completely frozen. OCT-embedded tissue
80 blocks were removed from the Cryomold and mounted on the specimen stage. 10µm sections were cut in
81 a cryostat at -10°C and placed within a Capture Area on the pre-equilibrated Visium Spatial Slides (10x
82 Genomics). The slides were later sealed in individual 50 ml Falcon at -80°C ready for further processing.

83 10X Genomics Visium experiments

84 Four short axis sections of the heart were processed according to the manufacturer's protocol. Libraries
85 were prepared individually, one per heart section. The Visium Spatial Tissue Optimization Slide & Reagent
86 kit (10x Genomics) was used to optimize permeabilization conditions. Tissue permeabilization was
87 performed for 24 min. Spatially barcoded full-length cDNA was generated using the Visium Spatial Gene
88 Expression Slide & Reagent kit (10x Genomics). A fraction of each cDNA library was used for Nanopore
89 sequencing. cDNA amplification was then conducted for 20 cycles of PCR (identified by qPCR), and
90 400µl were used in the 10xGenomics Visium library preparation (100µl per section). The libraries were
91 sequenced on a NovaSeq6000 (Illumina), with 29 bases from read 1 and 90 bases from read 2, at a depth
92 of 160M reads per section (640M reads in total). The raw sequencing data was processed with the 10x
93 Genomics Space Ranger v1.1.2 and mapped to the mm10 genome assembly (mm10-2020-A).

94 Oxford Nanopore sequencing libraries

95 Libraries for Nanopore sequencing were prepared according to the manufacturer's protocol for direct
96 cDNA Sequencing (SQK-DCS109 Oxford Nanopore Technologies) with the following minor modifications:
97 The protocol was started with 200ng cDNA with the End-prep step. Final library elution was performed in
98 15µl to have material left for TapeStation analysis and Qubit BR measurement. Four GridIon flow cells

99 (FLO-MIN106) were loaded with 12 μ L libraries (74-115ng) by using a Flow Cell Priming kit EXP-FLP002.
100 Base calling was done with Guppy v5.0.12. The High accuracy (HAC) model was selected for base calling
101 (Q-Score cut-off > 9).

102 Spatial barcode assignment

103 To account for source-specific quality differences, each heart section (Illumina libraries) was processed
104 separately using Scanpy v1.7.2 (Wolf et al., 2018), keeping only spatial barcodes with approximately 150
105 < counts < 18000, 250 < genes < 5000, detected in at least 2 spots, and with less than 40% mitochondrial
106 counts. The resulting datasets were concatenated, normalized, and the union of highly variable genes (per
107 section) were kept for final analysis. Batch balanced KNN (BBKNN) (Polański et al., 2020) with ridge
108 regression (Park et al., 2020) was used for integration and batch correction, starting from a coarse clustering
109 obtained from a BBKNN-corrected graph.

110 For the Nanopore libraries, samples were demultiplexed and processed with ScNapBar v1.1.0 (<https://github.com/dieterich-lab/single-cell-nanopore>) using a naive Bayes model to
111 assign spatial barcodes (Wang et al., 2021). Briefly, for each heart section, the Space Ranger results
112 (Illumina libraries) were used to parametrize a model of barcode alignment features to discriminate correct
113 versus false barcode assignment in the Nanopore data. FASTQ files were mapped using minimap2 v2.21 (Li,
114 2018). For transcript isoform quantification, a *de novo* transcriptome annotation was generated. Alignment
115 files were processed by StringTie v2.1.5 in long read mode with the reference annotation to guide the
116 assembly process (Kovaka et al., 2019). The annotations were merge into a non-redundant set of transcripts
117 and compared to the reference using GffCompare v0.12.2 (Pertea and Pertea, 2020), after removing single-
118 exon transcripts. To generate feature-spatial barcode matrices, alignment files were split into multiple files,
119 one per spatial barcode, based on the barcode assignments, converted to FASTQ, and aligned to the *de*
120 *novo* transcriptome with minimap2. Abundances were quantified with Salmon v1.5.2 in alignment-based
121 mode using a long read error model (Patro et al., 2017). Each section was processed separately using
122 Scanpy v1.7.2 and integrated with BBKNN, as described above for the Illumina data. Spatial barcodes
123 were filtered for counts (approximately 50 < counts < 4000), transcripts (approximately 50 < transcripts
124 < 2000, detected in at least 2 spots), and ribosomal genes (< 40%). Neighbors enrichment and cluster
125 co-occurrence analyses were performed using Squidpy (Palla et al., 2021).

127 Identification of anatomical regions

128 For the Illumina libraries, the neighborhood graph was computed using BBKNN. Spots were clustered
129 with a low resolution (0.3) to identify anatomical regions such as infarct, border and remote zones. Marker
130 genes were identified using a Wilcoxon rank sum test with Benjamini-Hochberg correction, by comparing
131 the expression of each gene in a given cluster against the rest of the spots. The final clusters were manually
132 annotated.

133 Labels were then assigned to Nanopore spatial barcodes based on the set of matching Illumina barcodes.
134 However, not all assigned barcodes were labeled due to quality control filtering criteria that were different
135 between Illumina and Nanopore datasets. To assign labels to the remaining Nanopore barcodes, seed
136 labelling was performed with scANVI using the set of assigned labels as groundtruth (Xu et al., 2021). The
137 top expressed transcripts were then identified as described above for the Illumina data.

138 Spatial spot deconvolution

139 Spatial spots were deconvoluted using stereoscope (scvi-tools v0.15.0) (Andersson et al., 2020). Heart
140 data (Smart-Seq2 and 10x Genomics) from the Tabula Muris (Schaum et al., 2018) were used as reference
141 dataset and highly variable genes were identified. For the Smart-Seq2 data, gene length normalization was
142 applied. The model was trained on the single cell reference dataset on the intersection of genes found in
143 the spatial (Illumina) data, and proportions were inferred for each Visium spot for each cell type in the
144 reference dataset. Labels were then assigned to Nanopore spots are described above.

145 Spatial gene expression (Illumina)

146 For each heart section, genes with spatial expression patterns were obtained with SPARK (FDR <
147 0.05) (Sun et al., 2020). Overrepresented biological processes in each region were identified using spatially
148 variable markers that were previously identified ($\log FC < 0.2$, p-value < 0.05), using Enrichr with the
149 Python package GSEApY.

150 Differential transcript usage (Nanopore)

151 To identify changes in relative usage of transcripts/isoforms within genes, differential transcript usage
152 (DTU) tests were performed using satuRn (Gillis et al., 2021). Only multi-exon transcripts and genes
153 with more than one isoform were kept for the analyses. The transcript count matrix was further filtered
154 to keep transcripts expressed in a worthwhile number of spots, determined by the design (but greater
155 than at least 10% of the smallest group size), with a CPM count above a threshold of 1(median library
156 size) $^{-1}$. In addition, transcripts were kept only if they had a minimum count of 1 across all spots. A quasi-
157 binomial generalized linear model was fit using a design comparing each anatomical region with another,
158 or each anatomical region with the rest of all regions. A two-stage testing procedure was performed using
159 stageR (Van den Berge and Clement, 2021), with an OFDR of 0.05. Results were reported using a student's
160 t-test statistic, computed with estimated log-odds ratios. To identify isoform-switching genes, significant
161 genes were identified for each contrast with at least two transcripts showing a switching expression pattern
162 between regions of interest.

163 Transcript classes

164 Transcript classes were assigned in scNAST using Gffcompare (Pertea and Pertea, 2020). For the
165 identification of isoforms and transcript classes between regions, markers previously identified were used to
166 select transcripts in each region with a $\log FC > 0.1$ (p-value < 0.05). Enrichment of transcripts of a certain
167 class (e.g. intron retention) was calculated using a Fisher exact test, using only non-equality classes. Motif
168 enrichment was performed using Simple Enrichment Analysis (SEA) from the MEME suite, using retained
169 intron sequences vs. annotated (reference transcript) sequences (Bailey et al., 2015).

RESULTS

170 scNAST enables the demarcation of spatially distinct regions of the myocardium post-MI

171 Fresh-frozen tissue samples were stained, imaged and fixed on Visium Spatial Gene Expression Slides
172 (10X Genomics) for permeabilization and *in situ* RNA capture. Full-length cDNA libraries were split for
173 the preparation of 3' Illumina short-read and direct long-read Nanopore sequencing libraries. Short-read
174 data were used for the assignment of spatial barcodes to Nanopore reads using the scSNAPBAR workflow,
175 and subsequently used to define anatomical regions within the tissue organization (Fig. 1A). Long-read

176 data were used for transcriptome assembly and transcript abundance quantification, and layered onto the
177 stained images to reveal the spatial organization of isoform expression. The Nanopore data comprises
178 of four heart slices (or samples) with a total of 25,5 million reads, reaching a relatively high sequencing
179 saturation (Fig. 1B), and providing a significant gain in coverage along full-length transcripts (Fig. 1C).
180 After spatial barcode assignment, libraries had a median of 2.8 million reads per sample, with over 70%
181 assigned reads (Figs. 1D and Supplementary Figure S1A). Despite variations across samples, we observed
182 a good per spot correlation between Illumina and Nanopore libraries (Fig. Supplementary Figure S1B,C).
183 Per spatial spot ($55\mu\text{m}$), after quality filtering, we observed a median read count varying between 593 and
184 2021 corresponding to a median of 311 (respectively 1000) distincts isoforms (Fig. Supplementary Figure
185 S2, see also Figs. Supplementary Figure S3 Supplementary Figure S4).

186 Clustering based on short-read gene expression defined four broad morphological regions: two remote
187 zones that stemmed from differences in sequencing depth between heart slices, a border zone, and the infarct.
188 Remote zones and, to a lesser extent the border zone, were largely associated with cardiomyocyte markers,
189 while the border and infarct zones were characterized by a higher expression of endothelial, myofibroblast,
190 and immune marker genes, as well as with markers of fibrosis and inflammation (Fig. 2A). The region
191 classification was then transferred to the Nanopore data (Fig. 2B), and markers of each region were identified
192 (Fig. Supplementary Figure S5). We also investigated, using the Illumina libraries, whether our spatial
193 transcriptomics data reflected known biological processes of myocardial infarction. We identified spatially
194 variable genes across samples and characterized each region using biological processes (Fig. Supplementary
195 Figure S6, and supporting information **Supplementary Table S1**). Remote zones were generally associated
196 with cardiac muscle contraction linked to an overrepresentation of healthy cardiomyocytes, and included
197 genes such as *Strt1* (SERCA regulator DWORF), cardiac troponin I (*Tnni3*), myosin-binding protein C
198 (*Mybpc3*), or ventricular myosin light chain-2 (*Myl2*). The border zone was characterized by genes such
199 as *Nppb* or *Ankrd1*, both of which were reported to be upregulated in the border zone after MI (Hama
200 et al., 1995; Mikhailov and Torrado, 2008), and other genes associated with the complement system and
201 myogenesis. Overrepresented processes were however similar to those found in the infarct area. Top
202 markers of the infarct area included collagens and genes associated with TGF- β and p53 signalling, with
203 hypoxia, coagulation, epithelial to mesenchymal transition, and the extracellular matrix. Overrepresented
204 processes included neutrophil degranulation, and gene expression associated with the innate immune
205 system. Remote, border, and infarct zones had a clear distinct spatial organization: the two remote zones
206 co-occurred at short distances with one another, but did not show any neighborhood enrichment with the
207 border and infarct zones (Fig. 2C,D). We observed a slight co-enrichment of the infarct and the border
208 zone at medium distances (Fig. 2D), revealing how spatial resolution may, in general, affect morphological
209 classification.

210 Following this strategy, we successfully assigned 7,616 spatial spots, corresponding to distinct 19,794
211 transcript isoforms in total, encoded by 12,474 genes. Among all genes, we observed 8,131 (67,3%) that
212 expressed a single isoform and 3,953 (32.7%) that expressed 2 or more isoforms (Fig. 2E). Although
213 predicted by our assembly, genes with many isoforms were expressed at a lower threshold and were
214 not included in our final analyses. We observed variations in the number of isoforms per gene across
215 morphological regions of each heart slice, with significant differences between either of the remote zones
216 and the border and the infarct areas, suggesting the existence of a higher transcriptome diversity in the
217 healthy regions (Fig. 2F).

218 scNAST reveals the spatial isoform diversity of the myocardium post-MI

219 Transcript isoforms were largely associated with exact matches to the reference annotation, multi-exon
220 transcripts with at least one junction match to the reference (*e.g.* exon skipping and exon extension),
221 transcripts longer than the reference (containment of reference), completely novel transcripts (intergenic),
222 transcripts with exonic overlap, intronic transcripts, or retained introns (Fig. 3A). Among these classes,
223 we were particularly interested in novel transcripts associated with retained introns. Intron retention (IR)
224 occurs when an intron is transcribed into pre-mRNA and remains in the final mRNA. Only recently has IR
225 become of interest due to its associations with complex diseases (Zhang et al., 2020). Interestingly, among
226 differentially expressed transcript markers, we found across the infarct and border zones 65 distinct IR
227 transcripts, compared to 1,612 in the remote zones, corresponding to an odds ratio of 1.73 (p-value 0.008).
228 When considering the infarct area only, the odds were even greater (odds ratio 2, p-value 0.002). Introns that
229 were retained across the infarct and border zone were enriched in motifs associated with poly(C)-binding
230 protein 1 and 3 (Pcbp1/3), KH RNA-binding domains (Khdrbs1), and Rbm38, a homolog of Rbm24, a
231 pivotal cardiac splice factor (Weeland et al., 2015) (supporting information [Supplementary Table S2](#)). The
232 genes harboring these transcripts were enriched in RNA metabolic/catabolic processes, RNA binding,
233 and the unfolded protein response (Fig. 3B). Among these, we found cardiac muscle alpha actin (Actc1),
234 the major protein of the thin filament responsible for cardiac contraction (Fig. 3C). Our data revealed
235 a relatively high expression of the IR transcript isoform across all regions, but a comparatively greater
236 contribution to the infarct area (Fig. 3D). Interestingly, Actc1 is located between two (antisense) long
237 non-coding RNAs, one of which has recently been described for its role in cardiomyocyte proliferation and
238 cardiac repair (Ponnusamy et al., 2019). At the Actc1 locus, we identified two novel antisense transcripts
239 with a good coverage across all regions (Fig. 3C). There is now growing evidence that antisense-mediated
240 gene regulation is involved in different pathophysiological contexts, including heart disease (Luther, 2005;
241 Zinad et al., 2017; Celik et al., 2019). Using the Tabula Muris (Schaum et al., 2018) as a reference dataset to
242 perform deconvolution, we identified cardiomyocytes, and to a lesser extent, endothelial and myofibroblasts
243 as the predominant origin of these novel antisense isoforms, with a correlation pattern that matched that of
244 the largest Actc1 annotated isoform (Fig. 3E).

245 scNAST allows to study regional isoform switching with spatial context

246 Among the multi-isoform genes, we looked for those showing a splicing pattern variation across
247 regions. We identified 109 significant regional isoform switching genes across all comparisons between
248 the remote, border, and infarct areas (OFDR 0.05 at both gene and transcript level, supporting
249 information [Supplementary Table S3](#)). Our results showed a clear regional isoform switching among
250 differentially used transcripts for genes involved in cardiac muscle contraction and tissue morphogenesis,
251 many of them clinically relevant, such as Ankrd1 (Mikhailov and Torrado, 2008), Sparc (McCurdy et al.,
252 2011), or Clu (Turkieh et al., 2019) (Fig. 4A). The largest number of isoform-switching genes were found
253 between the infarct and the remote zones (Fig. 4B). Among them, we found PDZ and LIM domain protein
254 5 (Pdlim5), a gene encoding for a protein that localizes to the Z-disk by binding to α -actinin, and which has
255 been implicated in dilated cardiomyopathy (Verdonschot et al., 2020) and in heart failure with preserved
256 ejection fraction (Soetkamp et al., 2021). Pdlim5 undergoes extensive splicing in both human and mouse.
257 Our data showed a significant change in isoform usage between two annotated protein-coding isoforms
258 between the infarct and the border or the remote zones (Fig. 4C,D,E). The shorter isoform was mostly
259 found in the border and remote zones, while the longer isoform was significantly, albeit at lower levels,
260 expressed in the infarct area (Fig. 4C,E).

DISCUSSION

261 SCNaST expands the spatial transcriptomics toolbox to third-generation long-read sequencing. SCNaST is
262 based on commercially available platforms, and can be used to investigate the isoform landscape of complex
263 tissue. We presented here a transcriptome-wide approach to explore annotated and *de novo* isoform diversity
264 with morphological context in the mouse heart after myocardial infarction (MI). Our approach opens-up
265 new opportunities to understand the spatial and molecular organization of the heart following injury. The
266 infarcted heart has usually been divided into three zones: the infarct, an ischemic border zone, roughly
267 defined as the area immediately adjacent to the infarct, and a remote zone. Although previous studies
268 showed transcriptional differences between proximal and distal areas to the infarct (van Duijvenboden
269 et al., 2019b), observations were limited to conventional short-read sequencing approaches or were limited
270 in spatial resolution. Our results not only reflected known biological processes of myocardial infarction,
271 but also revealed the spatial organization of gene expression and transcriptome diversity consistent with
272 the underlying biological condition. We identified novel isoforms in clinically relevant loci and regional
273 isoform switches that could play a role in the myocardium post-MI. We envision that our work will serve
274 as a reference for future developments and studies integrating long-read sequencing with spatial gene
275 expression data.

DATA AVAILABILITY

276 The datasets generated for this study can be found in the [to be deposited at SRA]. Additional
277 information, supporting data and scripts to generate the figures are available at <https://github.com/dieterich-lab/ScNaST>.
278

CODE AVAILABILITY

279 SCNAPBAR is publicly available at:
280 <https://github.com/dieterich-lab/single-cell-nanopore>. SCNaST is publicly
281 available at:
282 <https://github.com/dieterich-lab/ScNaST>.

AUTHOR CONTRIBUTIONS

283 CD, FL supervised the research and acquired funding. JA planned and supervised Visium experiments
284 and library preparation. XL performed MI experiments and tissue extraction. CB, RC performed Visium
285 experiments and library preparation. EB was in charge of software development, analyzed the data and
286 wrote the manuscript.

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CONFLICT OF INTEREST

292 The authors declare that they have no conflict of interest.

SUPPLEMENTAL DATA

293 **Supplementary Figure S1**

294 scNAST methodology.

295 **Supplementary Figure S2**

296 Quality control (Nanopore).

297 **Supplementary Figure S3**

298 Quality control (Illumina).

299 **Supplementary Figure S4**

300 Spatial distribution of UMIs or counts and genes or transcripts.

301 **Supplementary Figure S5**

302 Top markers of each region.

303 **Supplementary Figure S6**

304 Enriched processes among spatially variable marker genes.

305 **Supplementary Table S1**

306 Spatially variable genes.

307 **Supplementary Table S2**

308 Motif enriched in IR transcripts.

309 **Supplementary Table S3**

310 Differential transcript usage (isoform-switching genes).

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FIGURE CAPTIONS

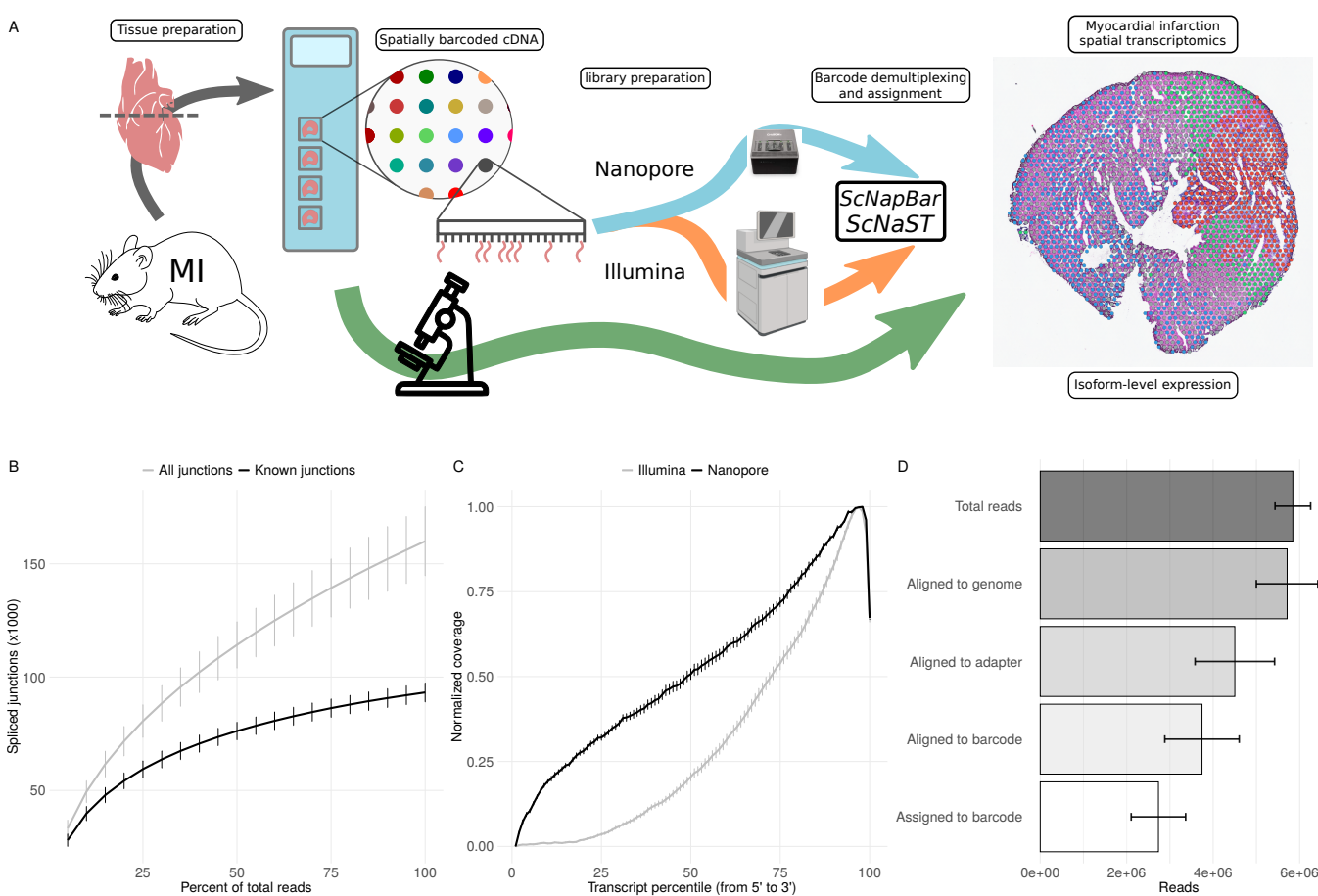


Figure 1. scNAST methodology. A Schematic of the scNAST workflow using a hybrid sequencing approach on Nanopore and Illumina platforms to assign spatial barcodes to long-read sequencing. B Nanopore sequencing saturation showing the number of splice sites detected at various levels of subsampling. A curve that reaches a plateau before getting to 100% data suggest that all known junctions in the library have been detected. The curve shows the mean \pm SE of four samples. C Normalized transcript coverage for Nanopore and Illumina. The curves show the mean \pm SE of four samples. D Reads assigned by SCNAPBAR at each step of the workflow. The bars show the mean \pm SE of four samples.

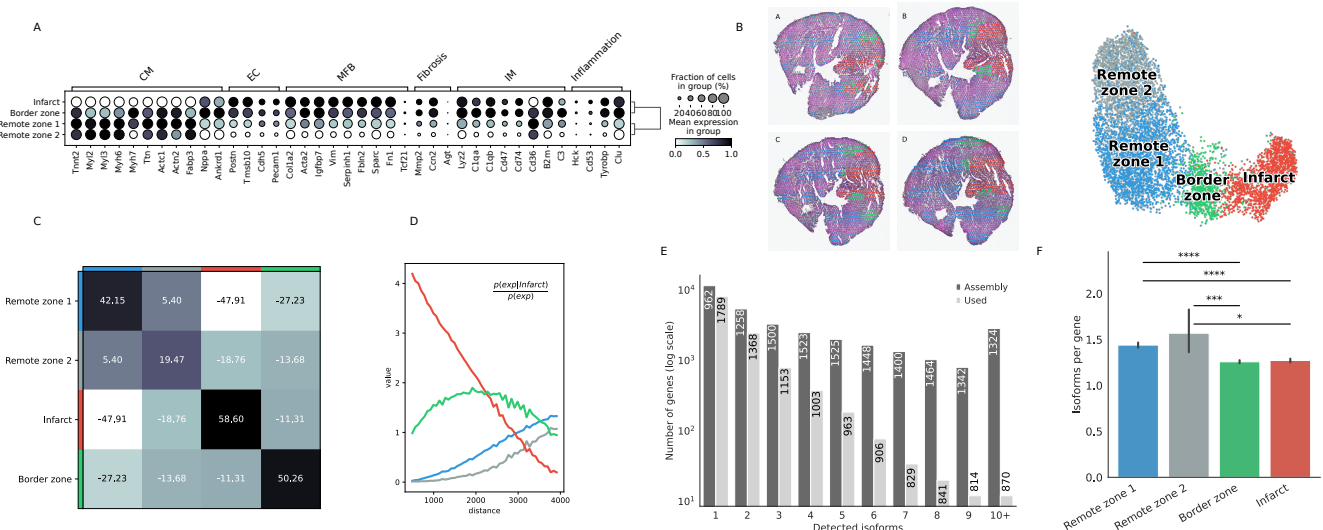


Figure 2. Defining morphological regions after MI. **A** Dot plot showing the expression of selected markers associated with the expression of CM=cardiomyocytes, EC=endothelial cells, MFB=myofibroblasts, IM=immune cells, or with fibrosis and inflammation, based on the short-read Illumina data. **B** Annotation of mouse heart regions after MI via short-read clustering, transferred to the Nanopore data. Scatter plot in spatial coordinates of the anatomical regions (left) and UMAP representation of the Nanopore data using the region annotation from short-read clustering (right). Colors in the spatial scatter plot are matching those of the UMAP. **C** Neighbors enrichment analysis in one heart axis section. The heatmap shows the enrichment score on spatial proximity between the different anatomical regions. Spots belonging to two different regions that are close together will have a high score, and vice-versa. **D** Cluster co-occurrence in spatial dimensions in one heart axis section. Line plot showing the the conditional probability of observing a given region conditioned on the presence of the infarct region, computed across increasing radii size around each spots. Distance units are given in pixels of the Visium source image. **E** Barplot showing the frequency distribution of the number of isoforms per gene, either stemming from the assembly, or found in the final data after quality control filtering. The median length of transcripts is indicated in each bar for each category. **F** Average number of isoform per gene detected for each morphological region. Significance was measured using a Mann-Whitney U-test (** = <0.001, *** = <0.0001)

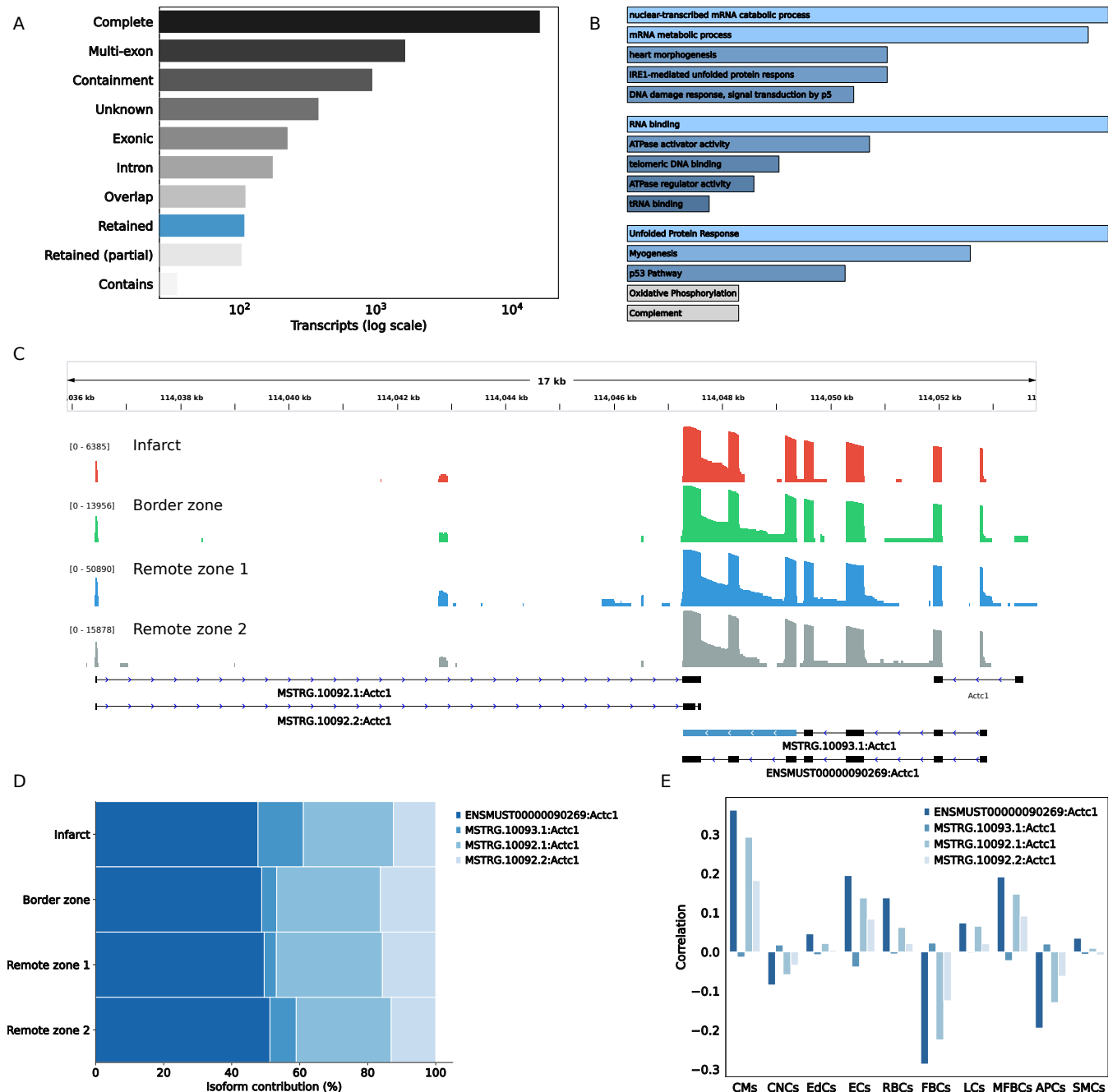


Figure 3. Characterizing the isoform diversity after MI. A Barplot showing how full-length transcripts obtained with scNAST compare to the existing mouse annotation. Labels Complete (=), Multi-exon (j), Containment (k), Unknown (u), Exonic (x), Intron(i), Overlap (o), Retained (m, n), and Contains (y) are explained in <https://ccb.jhu.edu/software/stringtie/gffcompare.shtml>. B Over-representation analysis of genes harboring novel transcripts with intron retention (IR). From top to bottom, biological processes, molecular function, and hallmark gene sets from the Molecular Signatures Database (MSigDB). C Exonic structure of the different *Actc1* isoforms, including novel isoforms identified by scNAST with intron retention and exonic antisense overlap. Coverage (log scale) is shown for each region with a different scale. D *Actc1* isoform contributions to total *Actc1* expression in the different heart regions. E Per spot correlation observed between spatial deconvolution of cell types and *Actc1* isoforms.

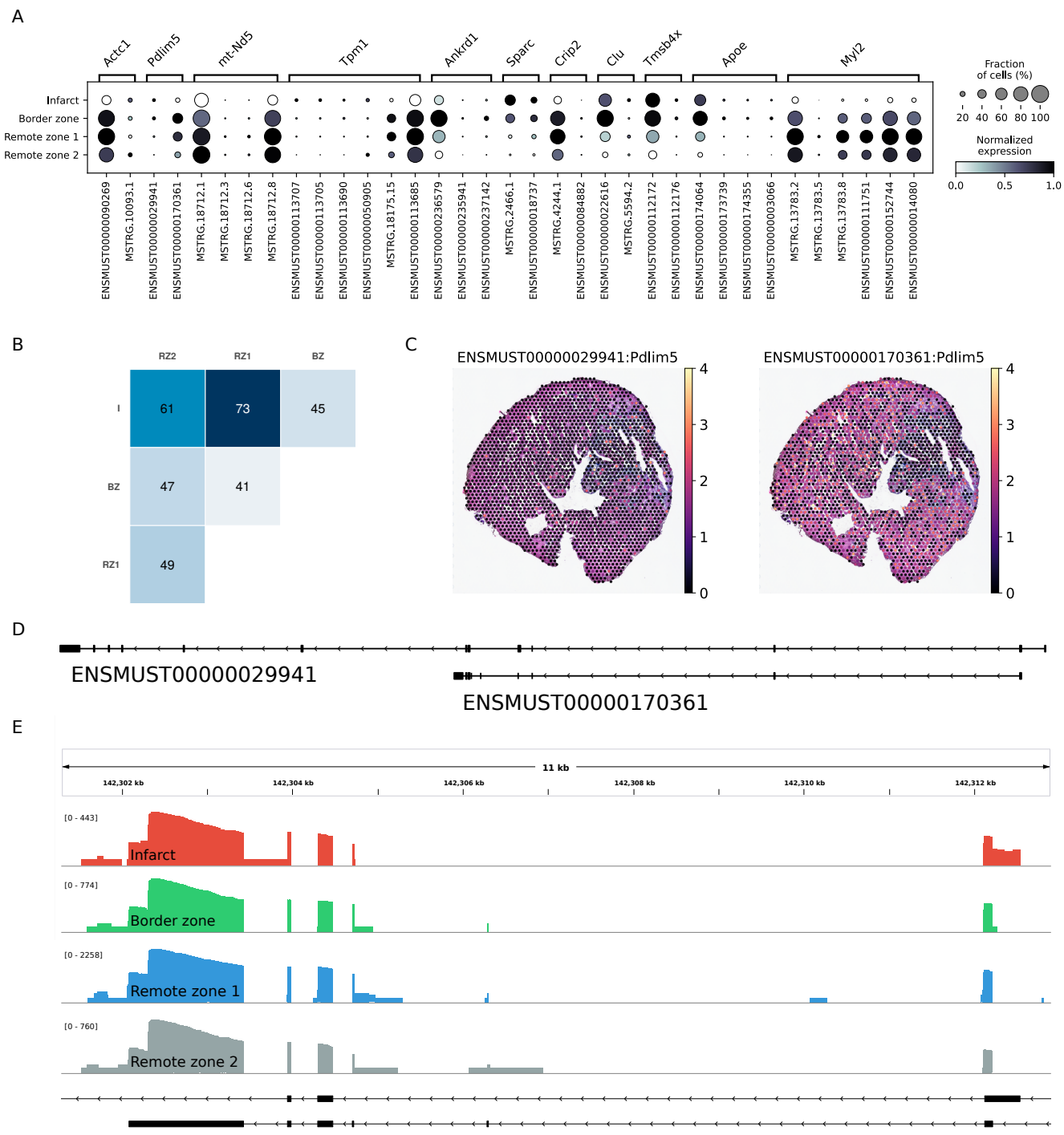


Figure 4. Regional isoform switching after MI. A Dot plot showing the top 5 most significant isoform-switching genes across all comparisons between the remote, border, and infarct areas using a stage-wise testing procedure at an overall false discovery rate (OFDR) of 0.05. The top genes were restricted to those identified as spatially variable. B Heatmap representing the number of isoform-switching genes identified between any two regions. C Pdlim5 isoform expression in one heart axis section. D Pdlim5 isoforms track. E Zoom of D to show coverage of both isoforms across regions.