Genomic and physiological analyses of the zebrafish atrioventricular canal reveal molecular building blocks of the secondary pacemaker region

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Abstract
The atrioventricular canal (AVC) is the site where key structures responsible for functional division between heart regions are established, most importantly, the atrioventricular (AV) conduction system and cardiac valves. To elucidate the mechanism underlying AVC development and function, we utilized transgenic zebrafish line sqet31Et expressing EGFP in the AVC to isolate this cell population and profile its transcriptome at 48 and 72 hpf. The zebrafish AVC transcriptome exhibits hallmarks of mammalian AV node, including the expression of genes implicated in its development and those encoding connexins forming low conductance gap junctions. Transcriptome analysis uncovered protein-coding and noncoding transcripts enriched in AVC, which have not been previously associated with this structure, as well as dynamic expression of epithelial-to-mesenchymal transition markers and components of TGF-β, Notch, and Wnt signaling pathways likely reflecting ongoing AVC and valve development. Using transgenic line Tg(myl7:mermaid) encoding voltage-sensitive fluorescent protein, we show that abolishing the pacemaker-containing sinoatrial ring (SAR) through Isl1 loss of function resulted in spontaneous activation in the AVC region, suggesting that it possesses inherent automaticity although insufficient to replace the SAR. The SAR and AVC transcriptomes express partially overlapping species of ion channels and gap junction proteins, reflecting their distinct roles. Besides identifying conserved aspects between zebrafish and mammalian conduction systems, our results established molecular hallmarks of the developing AVC which underlies its role in structural and electrophysiological separation between heart chambers. This data constitutes a valuable resource for studying AVC development and function, and identification of novel candidate genes implicated in these processes.

Keywords Atrioventricular canal · Atrioventricular node · Cardiac valve · Cardiac pacemaker · Zebrafish · RNA-seq
Introduction

The atroventricular canal (AVC) gives rise to the AV node, which constitutes part of the cardiac conduction system (CCS) responsible for generating and transmitting electrical impulses necessary for coordinated heart contraction [1, 2]. In the mammalian heart, the AV node can be found within the interatrial septum, at the AV junction [3]. Electrical impulses originating from the sinoatrial (SA) node are delayed by a fraction of a second in the AV node before being propagated further, ensuring consecutive contractions of the atrium and ventricle [4]. The AV node is often referred to as a secondary pacemaker as it possesses intrinsic automaticity, rendering it a potential arrhythmogenic source referred to as a secondary pacemaker as it possesses intrinsic automaticity [5, 6]. In the mammalian heart, the AV node can be found within the interatrial septum, at the AV junction [3]. Electrical impulses originating from the sinoatrial (SA) node are delayed by a fraction of a second in the AV node before being propagated further, ensuring consecutive contractions of the atrium and ventricle [4]. The AV node is often referred to as a secondary pacemaker as it possesses intrinsic automaticity, rendering it a potential arrhythmogenic source referred to as a secondary pacemaker as it possesses intrinsic automaticity [5, 6].

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of the heart [36, 38]. The sqet31Et transgenic line expresses EGFP in a ring structure marking the AVC [38], which likely corresponds to slow conducting myocytes homologous to the mammalian AV node [13, 14, 38, 39]. We utilize the in vivo labeling of the AVC in sqet31Et to isolate cells making up this structure and perform detailed molecular characterization by transcriptome profiling at 48 hpf and 72 hpf, corresponding to the time of CCS and cardiac valve development. To better understand the physiology of the CCS in zebrafish, we characterized electrical conduction patterns between the SAR and AVC, and cross-compared the transcriptome profiles of both pacemaker regions. We show that the AVC gene expression profile exhibits hallmarks of the mammalian AV node and reflects ongoing biological processes implicated in valve development. The transgenic line sqet33mi59BEt, in which the enhancer trap was inserted close by the fhf2 gene locus, expresses EGFP in the SAR [39]. Recently, we completed the analysis of the transcriptome of these cells [40]. Comparisons between the SAR and AVC transcriptomes revealed differences reflected in expression profiles of ion channels and connexins implicated in pacemaker function. A large number of AVC-enriched genes identified in this screen had human orthologs implicated in heart conditions related to cardiac conduction and valve/septal defects, suggesting the value of our transcriptome resource in identifying targets for further clinical investigations.

Results

Transgenic zebrafish line sqet31Et expresses EGFP in the zebrafish AVC

The sqet31Et transgenic line carries the Tol2 transposon containing EGFP gene driven by a 460-bp basic promoter from the zebrafish krt4 gene inserted in repetitive DNA region, which hinders mapping of the insertion site. The generation and detailed characterization of the line expressing EGFP in the bulbus arteriosus (BA) and AVC at high level and some myocardial cells at low level were described previously [38, 41]. To better visualize the GFP-expressing structure in sqet31Et, we performed high resolution imaging at 48 hpf and 72 hpf (Fig. 1A–D). We crossed sqet31Et with Tg(myl7:mRFP) that expresses membrane-bound RFP.
(mRFP) in cardiomyocytes. Confocal imaging of the AVC region revealed that at the surface of the AVC, the EGFP and mRFP expression overlapped, confirming the myocardial nature of the EGFP-expressing cells (Fig. 1C). At 72 hpf, two additional groups of ~3 cuboidal-shaped cells were detected at the deeper layer facing the cardiac lumen (Fig. 1D). These cells appear to be a part of the characteristic protrusion into the cardiac lumen most likely representing the developing AV cushion. Based on their location between the endocardium and myocardium, these cells have been previously defined as constituents of the non-chamber valve tissue [38, 39]. An additional EGFP expression domain detected previously [38] was observed at the BA from 72 hpf (Fig. 1B). The EGFP expressing domain in the developing heart of the sqet31Et transgenic line thus consists largely of AVC myocardium, with another expression domain in the BA, the latter observed only at 72 hpf.

**Transcriptome profile of the AVC**

To characterize the molecular profile of the GFP+ cell population in sqet31Et, we isolated these cells using fluorescence-activated cell sorting (FACS) at 48 hpf and 72 hpf (Fig. 2A) and profiled their transcriptome by RNA-seq. The rest of the heart cells, which did not express EGFP, were also collected (GFP−). Average sequencing reads mapping to the egfp sequence were considerably higher in GFP+ compared to GFP− samples, confirming the high representation of the EGFP-expressing cell population in the GFP+ samples (S1 Figure B). Principal component analysis (PCA) revealed tight clustering of replicates and clear separation between samples of different developmental stages (Fig. 2B).

To identify genes that are enriched in the AVC compared to the rest of the heart, we performed differential expression analysis between the GFP+ and GFP− fractions. In both developmental stages, a total of 3798 and 2777 genes were differentially expressed at 48 hpf and 72 hpf, respectively (absolute log2FC > 2, padj < 0.05), of which 1492 were common for both stages (Fig. 2C, F, S2 Table). GO and KEGG pathway enrichment analyses at 48 hpf revealed that the set of genes overexpressed in GFP+ compared to GFP− cells (absolute log2FC > 2, padj < 0.05; “AVC-enriched genes”) was overrepresented for functional terms related to cardiac muscle development and function (“cardiac muscle contraction”, “adrenergic signaling in cardiomyocytes”, “cardiac muscle development”, “cardiac muscle differentiation”, and “calcium signaling pathway”), in line with the myocardial identity of the GFP+ fraction (Fig. 2D, E, S3 Table). On the other hand, functional terms related to cell–cell adhesion (“cell adhesion molecules”, “cell–cell adhesion”) were overrepresented among transcripts overexpressed in the GFP− cell population. At 72 hpf, similar functional terms were overrepresented with the addition of “vascular smooth muscle contraction” term (Fig. 2G, H, S3 Table), which likely corresponded to the initiation of EGFP expression in the BA at this stage.

To assess whether GFP+ fraction contained AVC cells, we explored the presence of known markers of AVC in our dataset. We established a set of AVC marker genes for zebrafish and mammals by retrieving genes annotated with the term “atrioventricular canal” from the ZFIN (http://zfin.org/) and MGI [42] gene expression databases. Intersection of these known markers with the AVC-enriched transcriptome returned 46 and 58 genes in common, which were enriched in GFP+ cells at both 48 hpf and 72 hpf, respectively (Fig. 3A, B; S4 Table). Notably, several of these genes were known to be expressed specifically in zebrafish AVC myocardium, including bmp4 [24], wnt2bb [43], snai1b [44], hey2 [45], and hnf1ba [13]. On the other hand, 27 genes which were enriched in the GFP− population overlapped known AVC markers at either or both developmental stages. These included id4 and notch1b reported to be expressed in the endocardium [29, 46], which suggests that the GFP+ cells in sqet31Et are less likely to be endocardial.

Besides protein-coding genes, we found 108 and 19 transcripts defined as long intergenic noncoding RNAs (lincRNAs) according to the ENSEMBL database which were overexpressed in GFP+ cells at 48 hpf and 72 hpf stages, respectively (padj < 0.05; S5 Table). Among the lincRNAs overexpressed at 48hpf is the lincRNA ALIEN (linc.alien, ENSDARG00000097567) and si:ch211-265g21.1. The mammalian ortholog of linc.alien is known to be expressed in cardiovascular progenitors, and its function in cardiac lineage specification was demonstrated in both mammals and zebrafish [47]. The lncRNA si:ch211-265g21.1 was previously reported to be expressed in the embryonic zebrafish heart [48]. Additionally, MALAT-1 (malat1) which was reported to be expressed in heart [49] was found to be differentially expressed between 72 and 48 hpf. The rest of the lincRNAs in our list were mostly uncharacterized, which suggests the value of our transcriptome resources for discovering novel factors contributing to AVC development.

Besides the known AVC markers, the AVC-enriched transcriptome consisted of many other transcripts which were not previously associated with AVC or heart processes. To further validate the AVC-enriched transcriptome, we performed WISH on eight selected AVC-enriched transcripts with no previous heart expression reported. All of these were expressed in the AVC except for one, si:dkey-57k2.6, which is expressed only in the BA (S2 Figure). Another transcript, si:dkey-164f24.2, was expressed in the whole heart including the AVC (S2 Figure). Therefore, utilizing the sqet31Et transgenic line to specifically enrich for the GFP+ cell population, our RNA-seq analysis revealed the transcriptome representing the AVC, with contribution from the BA at 72 hpf.
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**Fig. 2** Transcriptome profiling of the GFP+ cells isolated from sqe-t31Et. A Scheme of experimental design. B Principal component analysis (PCA) on normalized RNA-seq data (regularized log) showing variance between three technical replicates of each sample as well as between samples. C-E Volcano plots showing genes differentially expressed between EGF-positive and -negative cells at 48 hpf (C) and 72 hpf (F). DESeq2 was used to calculate log2FC and padj values. Green spots indicate genes considered as significant (padj < 0.05) with at least twofold change between groups (log2FC > 2). Accordingly, light red spots represent significant genes with log2FC smaller than -2. D, G KEGG pathway enrichment of differentially expressed genes at 48 hpf and 72 hpf. Enrichment analysis was performed on a gene list meeting the following criteria: log2FC > 2 or log2FC < 2 and padj < 0.05. Same criteria were used to perform biological process Gene Ontology terms enrichment on up- and down-regulated genes (E, H). Top ten terms for each enrichment analysis are shown.
Fig. 3  AVC gene signatures are enriched in sqet31Et EGFP-expressing cells. Signatures were retrieved from MGI and ZFIN databases and used to identify molecular markers associated with AVC expressed in the studied dataset (padj < 0.05). A Heatmap depicts the dynamic of changes of known molecular AVC signatures that are in common across the developmental stage. B AVC markers uniquely expressed in either 48 hpf or 72 hpf stage. C Expression (in log10TPM + 1)) of genes encoding connexins, components of gap junctions which confer conductance properties between cells, in GFP+ and GFP− cells at 48 hpf and 72 hpf. Mammalian homologs of each connexin gene and their known conductance properties are described in the accompanying table.

To further discriminate the BA-associated genes from the AVC in the 72 hpf dataset, we intersected our data with a previously reported BA transcriptome of the adult zebrafish [100]. We found 56 out of 59 known BA-expressed genes enriched in GFP+ cells at 72 hpf (S6 Table). Unlike the AVC myocardium, the BA is mainly composed of smooth muscle [50]. Accordingly, in the 72 hpf dataset, we observed the enrichment of transcripts encoding smooth muscle light chain kinase (mylka), elastin B (elnb) and elastin A (elna) which is known to promote the differentiation of smooth muscle cell [51], as well as ltbp3 and fbln5 implicated in maturation and function of elastin [51–53] (S6 Figure). Therefore, although the 72 hpf GFP+ transcriptome contained genes expressed in the BA in addition to AVC, their largely distinct tissue composition, as well as the availability of BA transcriptome data [54], allowed us to distinguish between them. On the other hand, the expression of EGFP in the AVC and BA of the sqet31Et transgenic line adds to a list of common markers of these cell lineages.

**AVC gene expression profile shows signatures of AV pacemaker**

A hallmark feature of mammalian AVC myocardium is slow conduction, a property conferred by the composition of gap junctions between cells [9, 10]. Among the transcripts enriched in the GFP+ cell population, we found ten encoding various connexins. These consisted of cx36.7 (ortholog of human CX31.9 and murine Cx30.2 [55, 56]) and cx43.4/cx44.2 (ortholog of human CX45 [56]), whose mammalian orthologs form low or ultralow conductance gap junctions in the AV node [11, 12, 57]. These were enriched in the GFP+ population at both developmental stages, while other connexin transcripts were enriched only at 48 hpf (padj < 0.05). The latter group consisted of cx28.9/cx32.3 (ortholog of human CX37 [56]), cx43, and gja3/cx46 (Fig. 3C; S7 Table). Among those with the highest fold change between the GFP+ and GFP− cell populations were cx36.7, gja3/cx46, and cx32.3 at 48 hpf and cx36.7 at 72 hpf. Loss of function of mammalian CX46 leads to cardiac conduction disorders, while the loss of gja3/cx46 in the zebrafish mutant dco causes defects in heart morphology and ventricular conduction pattern [58]. The enrichment of genes encoding connexins forming low conductance gap junctions likely reflects the conduction delaying property of the AVC region. Besides those known for their role in cardiac conduction, transcripts encoding other members of the connexin family [Cx30.3 (CX30), Cx34.4 (CX30.3) and Cx35.4 (CX31)] were also enriched in the GFP+ cell population. These have not been previously implicated in heart or pacemaker function and are candidates for further investigation.

Besides delaying electrical conduction between atrium and ventricle, the AV node also possess intrinsic pacemaker activity [3, 5]. To determine whether this feature is conserved in the zebrafish, we searched amongst the GFP+ gene list for those known to be expressed in the AV node or associated with pacemaker development and function (S8 Table; [9, 59, 60]). Confirming previous reports [21], hcn4 expression was observed in GFP+ cell population at both 48 hpf and 72 hpf (S3 Figure; S8 Table). In addition, genes encoding zebrafish orthologs of Tbx18, Shox2, and Tbx2/3 [61–63] were expressed in the GFP+ population (S3 Figure, S8 Table). In mammalian CCS, Tbx2/3 are known to repress the expression of the chamber-specific Cx40 [33, 61]. In agreement with this, gja5a/b (the zebrafish ortholog of CX40) was not AVC enriched. It has been shown that nkd2.5 is expressed in all myocardium, but slightly higher in the AV conduction system [59]. Similarly, nkd2.5 was enriched in GFP+ cells compared to GFP− at 48 hpf (S3 Figure, S8 Table). Taken together, the transcriptome of AVC myocardium reveals conserved features to that of the mammalian AV node in terms of expression of genes linked to slow conductivity, automaticity, and molecular mechanism for AV conduction system development.

**Defect of the primary SAR pacemaker reveals spontaneous activity of the AVC**

The expression of hcn4 and other AV node markers in the zebrafish AVC led us to question whether it possesses inherent pacemaking activity as does its mammalian counterpart. We utilized isl1 K88X mutant (isl1m29), which exhibits a defective SAR pacemaker function manifested as sinus pauses and bradycardia [15, 18]. Apart from lacking the expression of fhf2a (Fig. 4A, B), bmp4 (Fig. 4C, D), and hcn4 (Fig. 4E, F) in the sinus venosus, isl1tm29 was devoid of EGFP-positive cells at the SAR, but not the AVC, as shown by analysis of sqet33ms59BEl [39] (Fig. 4G, H). Hence, we confirmed that the reduced number of cardiomyocytes at the venous pole in isl1tm29 observed previously [18] resulted from the absence of the pacemaker cells containing SAR. It is worth noting that isl1tm29 is the only vertebrate mutant that shows a complete lack of pacemaker SAR cells.
Despite the complete absence of SAR pacemaker, Isl1-deficient hearts contract, albeit inefficiently and irregularly, with long pauses (Fig. 5A, B, Supplementary movies 1 and 2). This suggests the existence of alternative origins of automaticity that triggers the initiation of cardiac contractions. To investigate whether the AVC could generate electrical impulses independent of the SAR, the Isl1 antisense morpholino [64] was injected into 1–4 cell stage zebrafish embryos expressing the genetically encoded voltage-sensitive fluorescent protein Mermaid, Tg(myl7:mermaid). This allowed direct observation of cardiac electrical conduction patterns. Similar to isl1 mutants, about 66% of Isl1 morphants showed sinus pauses (6 of 9 morphants vs 4 of 6 mutants), and all showed bradycardia. The heart rate of Isl1 morphants was 80.2 ± 15 beats per minute (bpm) at 48 hpf (mean ± SEM, n = 6) and 141.2 ± 10.3 bpm at 72 hpf, (n = 6), which is significantly lower than that in controls (176 ± 5.7 bpm at 48 hpf, (n = 7) and 229 ± 6.6 bpm at 72 hpf, (n = 18)) (Fig. 5A). Increased variability in heartbeat duration was noted in the Isl1 morphants as well.

In control hearts, the excitation wave front traveled uniformly across the atrium from the SAR toward the AVC (Supplementary movie 1; Fig. 5C, D). In contrast, in Isl1 morphants, the origin of the atrial excitation wave, although still predominantly from the SAR region, was more disperse and less coordinated, presumably driven by secondary pacemakers and automaticity of atrial cardiomyocytes (Supplementary movie 2; Fig. 5C, D). During the sinus pause, several stationary centers of automaticity distributed from posterior to anterior were detected, with the SAR region showing the highest activity. Interestingly, the location associated with AVC became more active near the end of the atrial contraction (Supplementary movie 2, Fig. 5C, D, asterisk), when triggering the subsequent phase of the excitation in the ventricle. The activity of the AVC location became even more pronounced during the periods of sinus pause. The loss of coordinated excitation wave and conduction in Isl1 deficient embryos suggests that the SAR is the primary pacemaker required for coordination of excitation wave. However, coordinated ventricular excitation can be induced by the electrical activity of the AVC with lower inherent pacemaker rate, in particular when no wave of excitation from the SAR drives the heartbeat.

**Comparison between the AVC and SAR transcriptomes reflect distinct electrophysiological properties**

The differences in the inherent activation rates of the SAR and AVC regions of the zebrafish heart led us to question the molecular nature underlying their distinct properties. We compared the transcriptome of the AVC with that of the SAR [40] to identify differentially enriched genes. Intersection between the two transcriptomes obtained a total of 1516 AVC-unique, 701 SAR-unique, and 450 common genes (S9 Table). Interestingly, while hcn4 was expressed in both SAR and AVC, its expression was enriched compared to the rest of the heart in the SAR but not the AVC. This may reflect the role of SAR as the dominant pacemaker (S9 Table). Several other transcripts encoding various ion channels were enriched in both SAR and AVC, notably, the T-type calcium channel Cacna1g, which is necessary for mammalian pacemaker activity in both SA and AV nodes [65]. Genes enriched only in AVC include trpm4 encoding a Ca²⁺-activated nonselective cation channel [66], which is implicated in human progressive familial heart block type I.
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Fig. 5 Effects of morpholino knockdown of isil on electrical activity of the atria in zebrafish larvae. A Heart rate at 48 hpf (left) and 72 hpf (right) in WT control (grey) and isil morphant (brown) zebrafish, showing slowed heart rate in isil morphants (Wilcoxon rank sum test with continuity correction, asterisks indicate *p value* ≤ 0.01). B Videographic analysis of the heartbeat in WT control (top) and isil morphant (bottom) 48 hpf zebrafish, showing slowed heart rate and sinus pauses (during the period indicated by the arrow) in the isil morphant. C Sequence of video frames showing electrical activation of the atria in WT control (left) and isil morphant (right) 48 hpf Tg(myl7:mermaid) zebrafish, showing normal activation (from the sinoatrial node region [SA] to the atrioventricular [AV] junction) and sites of latest activation/repolarisation (indicated by red stars) in the WT control and abnormal activation in the isil morphant (sites of early/ectopic activation in the morphant zebrafish indicated by green stars). D Isochronal activation map of the atria in WT control (left) and isil morphant (right) 48 hpf zebrafish derived from the video represented in (C)
characterized by cardiac conduction blockage downstream of the AV node [67]. Another notable example is cacna1c, whose human ortholog is associated with the Wolff–Parkinson–White syndrome, a condition affecting the AV conduction system [68, 69]. Other genes, including kcnq1, kcnq4, and apt1b1a, possess human orthologs associated with the maintenance of QT interval [70–72].

Despite having some common properties, the SA node serves a primarily pacemaking function, while the AV node is mainly specialized to delay electrical propagation between the atrial and ventricular chambers [9]. Therefore, while both regions express partially overlapping, mostly low-conducting gap junction proteins [9, 73], the AV node is particularly enriched in Cx30.2 and Cx45 [33, 74]. Accordingly, cx36.7, the zebrafish paralog of Cx30.2, was enriched in the AVC but not the SAR (S9 Table). On the other hand, cx43.4, a paralog of Cx45, was enriched in both the SAR and AVC (S9 Table). Low electrical coupling is also a necessary property within the definitive pacemaker cells of the SA node to prevent inhibitory interference from the surrounding working myocardium, which is more hyperpolarized [9]. Collectively, the overall differences in ion channel, cell adhesion, and extracellular matrix composition enriched in the SAR and AVC likely underlie their distinct electrophysiological properties.

**Developmental signaling pathways dynamics suggests signaling pathways implicated in valve formation**

Besides hosting the AV node, the AVC is also the site where cardiac valve formation is initiated. We identified transcripts encoding genes involved in the EMT process enriched in the GFP+ cell population (Fig. 6A, B; S10 Table). Members of the TGF-β signaling pathway were enriched in GFP+ cells at both developmental stages (tgfb1a, smad1, smad6b, and smad9) or specific to either stage (tgfb3, tgfb2b, and smad6a) (Fig. 6C). This is in line with the observation in mammalian endocardial cushion formation, where various TGF-β ligands are expressed in different cell populations of the AVC [75]. Notch signaling activity in the AVC endocardium is necessary for inducing EMT [76, 77] and transcripts encoding its key components were enriched in GFP+ cells at both stages, while wt1ab, wt6b, wt5ab, and wt4 were enriched only at 48 hpf (Fig. 6C). Genes encoding Wnt signaling receptors fzdb, fz9ab, fz2, fz6, fzd6, fzd8a, and fzd10 were enriched in both or either stages (Fig. 6C), whereas dkka/b and dkk2 encoding Wnt antagonists were enriched at 48 hpf. To visualize Wnt signaling activity in the AVC region, we crossed the sqe-t31Et transgenic line with the Wnt reporter line (Fig. 6E). The identification of these factors would allow deeper understanding of the molecular networks governing AVC formation and function.

**AVC-enriched genes are associated with human congenital heart defects related to CCS, valves and septa**

We identified human orthologs of AVC-enriched genes and interrogated them for any association with clinical phenotypes related to ClinVar terms: “arrhythmia”, “AV block”, “long QT syndrome”, and “conduction”. Our analysis revealed a total of 91 and 60 unique genes associated with these four ClinVar terms at 48 hpf and 72 hpf stages, respectively (S14 Table). Specifically, disease conditions represented by these terms included general forms of cardiac arrhythmia such as atrial fibrillation, sick sinus syndrome, abnormal QT interval, and Brugada syndrome, as well as those conditions specifically associated with defects of the AV conduction system or downstream effects such as heart block [82], Wolff–Parkinson–White pattern [69], and supraventricular tachycardia [83]. The latter group
Fig. 6 The transcripts of genes involved in EMT and valve development are enriched in the AVC. A AVC enrichment (expressed in log2-fold change between GFP+ and GFP− cells, padj < 0.05) of genes known to regulate EMT at both 48 hpf and 72 hpf stages. B Overlap of EMT-regulating genes enriched in GFP+ cells at both stages. C, D Volcano plot showing enrichment of components of the TGF-β and Wnt signaling pathways in AVC at both 48 hpf and 72 hpf stages. E Whole mount in situ hybridization of several AVC-enriched genes of the TGF-β and Wnt pathway components.
included trpm4 and cacna1c, which were enriched in both SAR and AVC, as well as myhpc3, smyhc2, hrc, dspla, myh7l, zgc:86709, lmmn, snat1, and ttn.2.

As the AVC is also the site where endocardial cushion and valve develop, we expected to find associations between AVC-enriched genes and human valve and septal defects. We searched human orthologs of AVC-enriched genes for overlap with ClinVar terms containing “tricuspid valve”, “AV valve”, “mitral valve”, and “valve in general”. In total, 115 and 93 unique genes were associated with these terms at 48 hpf and 72 hpf, respectively (S15 Table). In addition, 66 and 55 unique genes were associated with the term “septal defect” at each respective stages (S16 Table). In the adult human heart, AV node is embedded into the interatrial septum [3]. Given that the endocardial cushions are involved in the formation of the AV valves and septa, defects of interatrial septum could be linked to defects in cardiac conduction. In fact, a number of genes were commonly associated with ClinVar terms “cardiac conduction” and “valve” (S16 Table). For example, tbx5a, whose human ortholog TBX5 causes the Holt–Oram syndrome characterized by congenital heart malformation due to variable atrial and ventricular septal defects as well as heart conduction defects [74, 84]. Another notable example is smyhc2, whose human ortholog MYH6 is associated with both atrial septal defect and sick sinus syndrome [85, 86]. Other examples include cacna1c, ttn.2, snat1, lmmn, dspla, and myhpc3. The overlap of a large number of AVC-enriched genes with human heart conditions related to CCS and valve/septal defects suggests our transcriptomics data as a valuable resource for studying these diseases.

Discussion

The AVC constitutes part of the CCS which serves as the site where the propagation of electrical impulses is delayed, allowing consecutive contraction of the atrium and ventricle. In addition, it is also the site where the heart valves develop. The study of the AVC is challenging due to the lack of specific molecular markers defining this region. Available data relied on methods based on histological sections [87], which lacks the ability to isolate specific cell types. Nevertheless, it is known that distinct structures of the AVC, such as the pacemaker cells and cardiac valve tissue, express unique combinations of marker genes which can be used to distinguish them. The transgenic line sqet31Et provides the necessary level of specificity, which allows the enrichment of AVC myocardial cells by FACS. A caveat of this approach remains in the fact that the EGFP expression pattern in the sqet31Et transgenic line is driven by the activity of a yet unknown enhancer [38], which prevented an accurate assessment of the homogeneity of cell populations expressing EGFP in this transgenic line and their precise identity. For now, it is challenging to identify this enhancer due to the insertion of the enhancer trap construct in genomic repeat regions. However, with the increasing availability of long read sequencing methods, it may be possible in the near future to map the insertion site and trace the identity of this enhancer. Nevertheless, the co-expression of EGFP in the AVC and BA and cell populations suggests a unifying regulatory principle governing the specification of different cell types spatiotemporally. Transcription factors such as Tbx3 are expressed in both AV node and outflow tract mesenchyme, suggesting a similarity of developmental mechanism [88]. This poses an interesting question on gene regulation by the regulatory element(s) driving the expression pattern in sqet31Et transgenic line.

Our transcriptome analyses revealed that the zebrafish AVC myocardium possesses hallmarks of the mammalian AV node. The AVC transcriptome is characterized by high expression of mRNA encoding low conductance connexins cx36.7 and cx43.4, as well as the T-type calcium channel cacna1g and pacemaker hyperpolarization-activated channel hcn4. All these factors are known to define the AV node and pacemaker activity in the mammalian heart [11, 57, 65, 89, 90]. The conserved features also extend to the expression of transcripts encoding the core pacemaker transcriptional network consisting of Tbx2a/2b/3/18 and Shox2 transcription factors.

The existence of the CCS in zebrafish has been supported by optogenetic studies [20, 22] with other evidence suggesting that the endocardium and hemodynamic stimulation play an important role in its development [39, 46, 91]. The SA pacemaker has been relatively better characterized, and it has been shown that its activity depends on Isl1 [15]. Using the sqet33mi59BEt transgenic line, we show that the loss of Isl1 abolishes the SAR harboring the primary pacemaker activity. Analysis of electrical conduction patterns in isl1 morphants revealed disorganized excitation, which still generally progressed from the SA to AV, but included sites of ectopic automaticity. This suggests that by driving primary pacemaker function, Isl1 acts to coordinate atrial activation. However, we cannot rule out the possibility that the isl1 knockdown did not abolish its function completely. In either case, the lack of an organized atrial activation pattern affects overall cardiac contraction, indicating that the coordinated signaling from the SAR and its propagation play a crucial role in coordinating heart contraction. The increase in AVC activation during the pause in heart rhythm in Isl1-deficient embryos demonstrated that it possesses inherent automaticity, enabling it to independently excite when a weakened signal from the primary SAR pacemaker is not sufficient to drive heart contractions. This corroborates previous observations in adult zebrafish heart of spontaneous electrical
activity at the AV region following surgical uncoupling of
the ventricle from the atrium [20]. Comparison of the tran-
scriptome profiles of the SAR and AVC, while revealing
common markers of pacemaker activity such as c Rules and
cxx34.4, also indicated differences that would affect elec-
trical properties, such as the enrichment of distinct types of
ion channel, gap junction, and extracellular matrix com-
ponents. This further supports that, despite their shared ability
to act as a pacemaker, the SAR and AVC performs different
functions.

Mammalian AV node consists not only of definitive pacemaker
cells, but also fibroblasts, macrophages, and ECM, which provide
electrical insulation around the AV node [2, 30]. Electrical impulses from AV node are further propa-
gated by the His/Purkinje fiber network, which link to the
thick myocardial tissue throughout the whole ventricle [31].
In contrast, the two-chambered heart in most fishes resembles
the mammalian embryonic heart tube, where electrical
current is propagated from one end to the other by means of
electrical coupling of cardiomyocytes without a specialized
CCS [2]. Hearts of ectothermic animals contain no insulat-
ing fibrous structure, although the slow conducting muscles
of the AVC is present [92]. Moreover, teleost hearts are not
known to possess any defined Purkinje fiber network, and
conduction function is served by the ventricular trabeculae,
which form myocardial continuity between AVC and apex
of the ventricle [93]. Therefore, it is reasonable to assume
that CCS function could be served by an equally simplified
structure, in which a subset of cardiomyocytes performs
pacemaking function and at the same time express additional
attributes, which enable it to slow down electrical propaga-
tion. Intriguingly, it was previously observed that cells of
the embryonic SAR send processes into the AVC, which
appeared as a network connecting the two structures [39]. It
is therefore tempting to speculate that a previously unchar-
acterized structure or cell type may exist in the zebrafish,
which facilitates fast conduction between the SAR and AVC.

Besides serving as part of the CCS, the AVC is also the
site where the heart valves and septa develop. Accordingly,
the AVC transcriptome was enriched for transcripts encod-
ing regulators of EMT, which is a hallmark of endocardial
cushion and valve development. In addition, components
of major signaling pathways, including Wnt, Notch, and
TGF-β, which were implicated in endocardial cushion and
valve development [28, 94], were differentially expressed
at 48 to 72 hpf. Canonical Wnt signaling is known to play
multiple roles in valve development, including regulation of
AVC maturation and establishment of its electrical proper-
ties upstream of Tbx3 [81]. In the adult zebrafish heart, a
compact group of Hcn4-positive cells is embedded within
the musculature of the AV valves [20]. The close association
between the valve tissues and pacemaker cells is reflected
in our transcriptome and adds to the heterogeneity of cell
types present within this region. Currently, bulk RNA-seq
approach does not allow us to distinguish between the vari-
ous cell subpopulations, or to clearly demarcate the con-
current developmental processes within the AVC region.
Analyses at the single-cell level in both embryonic and adult
zebrafish hearts are ongoing which is expected to reveal the
true cellular diversity of this structure and more accurately
characterize the CCS organization in zebrafish heart.

Conclusions

Collectively, our results establish that the zebrafish AVC
possesses molecular and physiological hallmarks of a sec-
ondary pacemaker, similar to that of the mammalian AV
node, in terms of automaticity, low conductance properties,
and conserved expression of developmental genes. The par-
tially overlapping expression profiles of genes encoding ion
channels and connexins likely underlies the distinct conduc-
tion functions between the SAR and AVC. In addition, the
dynamic expression of signaling pathways implicated in the
ongoing valve development illustrates the role of the AVC
in both electrophysiological as well as structural separation
between the heart chambers. The AVC transcriptome data
generated in this study will enrich our knowledge of molec-
ular factors, including novel candidate genes and noncod-
ing transcripts, implicated in cardiac conduction and valve
development.

Methods

Zebrafish

Wild-type, sqet31Et and sqet33mi59BEt enhancer trap
[38, 39], and other zebrafish lines used in this study:
Tg(myl7:mRFP) [95], Wnt reporter line Tg(7xTCF-Xla.
Siam:nlsmCherry) [80], were maintained in the zebrafish
facility of the International Institute of Molecular and
Cell Biology in Warsaw (license no. PL14656251) in
line with standard procedures and ethical guidelines.
Tg(myl7:mermaid) was generated by injection of a Tol2-
myl7-mermaid construct (kind gift of Yasushi Okamura),
together with transposase RNA, into one- to two-cell
stage AB zebrafish embryos, followed by screening for
fluorescence progeny. The isl1 K88X mutant (isl1 ka29) and
Tg(myl7:mermaid) were bred and maintained at the Hare-
field Heart Science Centre according to the Animals (Sci-
entific Procedures) Act 1986.Embryos were raised in egg
water at 28 °C, screened for a fluorescence signal in the heart
and staged at 48 hpf and 72 hpf based on established mor-
phological criteria [96]. The generation and characterization
Heart extraction and fluorescence-activated cell sorting (FACS)

To isolate the heart, embryos were anesthetized with Tricaine (0.16 mg/ml in egg water) and large-scale extraction was performed according to a previously published protocol, with minor adjustments [98]. GFP-expressing hearts were manually separated from remaining tissue under a fluorescence stereomicroscope and collected into 0.5 ml of EDM (L-15/10% FBS). Pools of 300–500 hearts were dissociated with Trypsin–EDTA solution (0.05%) as previously described [99]. A FACS Aria II cytometer (BD Biosciences, USA) was used to enrich GFP positive (fluorescent) and GFP negative (nonfluorescent) heart fractions. Gates for cell sorting were calibrated against dissociated hearts extracted from wild type zebrafish embryos at the respective developmental stages (48 hpf and 72 hpf). On average, FACS yielded 15–25% of GFP+ events of total singlet events (Supplementary Fig. 1A).

RNA extraction

To obtain high-quality total RNA, cells were sorted directly to 500 µl TRizol™ LS Reagent (Thermo Fisher Scientific, USA) followed by RNA purification and DNase I treatment by means of a Direct-zol™ kit (Zymo Research, USA) according to the manufacturer’s protocol. The Tapestation 2200 and High Sensitivity RNA ScreenTape assay (Agilent Technologies, USA) together with Quantus™ Fluorometer (Promega, USA) were used to assess quantity and quality of total RNA. The average RNA Integrity Number equivalent (RIN°) for samples used for downstream analysis was 8.7.

Library preparation and sequencing

To obtain sequencing libraries, a two-step approach was applied. First, cDNA carrying full-length transcript information was synthesized with SMART-Seq® v4 Ultra® Low Input RNA Kit for Sequencing (TuKaRa Bio, Japan), followed by Nextera XT DNA Library Preparation Kit (Illumina, USA) according to the manufacturer’s guidelines. As previously, Tapestation 2200 and dedicated High Sensitivity D5000 ScreenTape and High Sensitivity D1000 ScreenTape assays were used to validate final cDNA and sequencing libraries, respectively. Final libraries were quantified with KAPA Library Quantification Kit Illumina® Platforms (Kapa Biosystems, USA), followed by paired-end sequencing (2 × 75 bp) performed with Nextseq 500 (Illumina, USA). Libraries were sequenced in triplicate, where a single replicate consisted of GFP-positive and GFP-negative fractions for both developmental stages (48 hpf and 72 hpf), at an average depth of 47 million reads.

Analysis of sequencing data

FastQC tool v. 0.11.8 [100] was used to assess the quality of obtained raw RNA-seq reads. Minor adapters contaminations were removed by Cutadapt v. 1.17 [101] and RNA-seq reads were mapped to the zebrafish reference genome (GRCh11) using Salmon tool v. 0.9.1 [102] resulting in an average of 75% mappability rate (S1 Figure C). Sequencing reads were further analyzed in R programming language v. 3.5.2 [103], whereas differentially expressed genes were identified by the DESeq2 package [104]. Principal component analysis was performed on normalized reads counts transformed to the log2 scale by plotPCA function from the same package. ClusterProfiler v. 3.17.3 [105] was used to calculate the enrichment of both biological processes of Gene Ontology terms as well as KEGG pathways. The enrichGO and enrichKEGG functions were used with default pvalueCutoff and qvalueCutoff parameters. The ggplot2 package [106] was utilized for plots generation. Discovery of lincRNAs was performed using Ensembl GRCh11 primary assembly (v. 103) and ZFIN.

Confocal imaging

Embryos used for imaging were grown in egg water supplemented with 0.003% 1-phenyl 2-thiourea (PTU) at 24 hpf to prevent the formation of melanophores and pigmentation. Prior to imaging, embryos were anesthetized with 0.02% tricaine (MS-222; Sigma-Aldrich A5040), embedded in 1% low-melt agarose (Sigma, USA) in egg water, and mounted in a glass-bottom dish before imaging on an inverted confocal microscope (LSM800, Zeiss). Images were further processed with Imaris 8 software (Bitplane).

Optical mapping of atrial excitation

To visualize excitation in the embryonic heart, a transgenic zebrafish line expressing the FRET-based voltage-sensitive fluorescent protein Mermaid [107] specifically in myocardial cells Tg(myl7:mermaid) was used and optical mapping was performed as described previously, with minor adjustments [108]. Injection of morpholino against isl1 (5′-TTAATCTGC GTTACCTGATGTAGTC-3′) was performed as previously described [64]. Taking into account sample distributions, variances, and sample sizes we decided to apply a non-parametric test for both conditions (Wilcoxon rank sum test with continuity correction) which resulted in p-value = 0.002165 (at 2 dpf) and p-value = 0.001483 (at 3 dpf). Embryos were embedded in 1% low melting agarose on a 35 mm Petri dish.
and oriented ventral side up to the imaging plane. Embedded embryos were transferred to an imaging chamber (RC-29; Harvard Instruments, USA) with a heated platform (PH-6D; Harvard Instruments). The temperature was maintained at 28 °C by a temperature controller (TC-344B; Harvard Instruments). Images were obtained using an epifluorescence upright microscope (BX51WI; Olympus) and focusing module (BXFM; Olympus) with a 40X water immersion objective (LUMPLFLN 40XW; Olympus) and magnification changer (U-CA; Olympus). Fluorescence was excited using a blue light-emitting diode (CBT-90; Luminus, USA) passed through a 460 ± 5 nm bandpass filter (HQ460/10X; Chroma, USA). Fluorescence was collected with a 482 nm dichroic mirror (FF482-Di01; Semrock, USA). To obtain simultaneous images of FRET donor and acceptor signals, dichroic mirror (FF552-Di02; Semrock), and passed through either a 500 ± 30 nm (HQ500/60 m-2p; Chroma) or 600 ± 37.5 nm (HQ600/75 m; Chroma) bandpass filter. Filtered emission was projected to two halves of a 16-bit, 128 × 128, 24 mm² pixels, cooled electron multiplying charge-coupled device camera (Cascade: 128 +; Photometrics, USA) and collected at 52 Hz with a ~ 19 ms exposure time. Images were processed and analyzed using custom routines in MATLAB ('wiener2') with a 3 × 3 pixel window. The atrium was manually segmented and each pixel signal normalized through time. Activation time was measured as the point at which the rate of voltage upstroke was maximal.

Electrophysiology

Micropipettes for electrocardiograph (ECG) measurement on whole zebrafish larvae were prepared by pulling fire-polished borosilicate glass capillaries (World Precision Instruments) using the Flaming/brown micropipette puller P-1000 (Sutter Instrument). The zebrafish larvae were mounted (laterally) in 1% low melting agarose in a glass dish and submerged in external buffer: 1 x egg water (0.6 g/L sea salt in reverse osmosis purified water). The micropipette was filled with internal buffer (174 mM NaCl, 2.1 mM KCl, 1.2 mM MgSO4.7H2O, 1.8 mM Ca(NO3)2.4H2O, 15 mM HEPES, pH 7.2) and the tip was positioned right above the pericardial region of the zebrafish heart. The electrical signals from the zebrafish heart received were recorded by pCLAMP 10 software (Molecular Devices) after amplification via Multiclamp 700B amplifier (Molecular Devices) and digitization through Axon Digidata 1440A digitizer (Molecular Devices). Data were analysed with Clampfit 10 software (Molecular Devices).

Whole mount in situ hybridization

For antisense probes generation, total RNA from 72 hpf embryos was extracted and reverse transcribed into cDNA with SuperScript IV Reverse Transcriptase (Thermo Fisher Scientific, USA). Obtained cDNA was used as a template for PCR. Purified PCR products were used as a template for in vitro transcription from the T7 promoter. Primers used are listed in S1 Table or reported previously [39]. Whole mount in situ hybridization (WISH) was performed as previously described, with minor adjustments [109]. Zebrafish embryos were grown in egg water containing PTU and fixed overnight at desired developmental stage in 4% paraformaldehyde in 1 × PBS (PFA/PBS). After sequential washes with 1 × PBT (50 ml 1 × PBS + 250 µl 20% Tween-20), embryos were digested for either 30 min (48 hpf) or 50 min (72 hpf) with 10 µg/ml proteinase K (Roche), washed with 1 × PBT, and fixed again for 1 h. PFA/PBS solution was discarded, and embryos were pre-hybridized overnight at 68 °C in a hybridization buffer. Subsequently, diluted and denatured probes were added to the pre-hybridized embryos followed by overnight incubation (68 °C) in a water bath. Post-hybridization washes were performed in increasing concentration of 2xSSC in the hybridization buffer. To reduce nonspecific signal, commercial blocking reagent (Roche) was used. Signal was visualized by overnight incubation with 1:5000 anti-DIG-AP antibody (Roche) at 4 °C followed by washing and addition of NBT and BCIP staining solution. After the staining was fully developed, staining solution was washed away and embryos were fixed in 4% PFA in 1 × PBS. For whole mount in situ imaging, embryos were mounted in glycerol and imaged on Nikon SMZ25 microscope. For each probe, WISH experiments were performed on at least 20 embryos obtained from at least three different breeding pairs.

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Author contributions CW and VK conceived the project and designed the experiments. KAN, AS and CW performed experiments. KAN prepared libraries and did sequencing. TAQ and PKL were in charge of the electrophysiological experiments. KAN and ML did experimental validation by light and confocal imaging. LB and KP planned and coordinated flow cytometry/sorting experiments. KAN, MM, SM, and MP contributed to data analysis. LJ performed analysis of isl1 mutants. CW, VK, TB, PK, and KAN wrote the manuscript. All authors read and approved the final manuscript.
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Data availability All sequencing data have been deposited in the GEO database under accession GSE160107.

Declarations

Conflict of interest The authors declare that they have no competing interests.

Ethical approval According to EU Directive 2010/63/EU on the protection of animals used for scientific purposes, the earliest life stages of animals do not fall into the regulatory frameworks of animal experimentation. This study used only zebrafish embryos less than 120 hpf which therefore does not require consent from the Local Ethics Committee.

Consent for publication Not applicable.

Consent to participate Not applicable.

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