Regulation of the diversification of the nodal and chamber myocardium
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Regulation of the diversification of the nodal and chamber myocardium
Regulation of the diversification of the nodal and chamber myocardium
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On the front cover: Whole mount fluorescence microscopy image of transgenic mouse hearts. A modified bacterial artificial chromosome (Bac337-Egfp) with the enhanced green fluorescent protein (GFP) gene inserted into the ANF gene drives atrial expression in a healthy control heart (left side) and is activated in the ventricles during heart failure (right side). On the back: Whole mount fluorescence microscopy image of an E13.5 transgenic mouse embryo carrying Bac336-Egfp. Egfp expression is restricted to the heart. See related Chapter 2.
Regulation of the diversification of the nodal and chamber myocardium

ACADEMISCH PROEFSCHRIFT

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aan de Universiteit van Amsterdam
op gezag van de Rector Magnificus
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ten overstaan van een door de college voor promoties ingestelde commissie, in het openbaar te verdedigen in de Agnietenkapel
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door

Thomas Horsthus

geboren te Weerselo
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'Because it's there'

British mountaineer George H.L. Mallory (1886-1924) replied to the question 'Why do you want to climb Mt. Everest?'
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**Summary**  

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Scope

The heart is the first organ that is formed during development. In human it beats after 3 weeks of development, comparable with embryonic day E8.5 in mouse. At this early stage the heart is a contracting linear tube, of which the myocardial cells share important features, among which high automaticity, low conduction velocity and low contractility. By an interplay of proliferation, differentiation, cell growth and addition of extracardiac cells, the primitive tube eventually gives rise to the adult 4-chambered heart, containing many phenotypically different cells. When the atrial and ventricular chambers differentiate from the embryonic myocardium, they obtain the so-called ‘working’ myocardial phenotype, characterized by high contractility and conduction velocity, but low automaticity. The nodal components of the conduction system of the adult heart, the sinus node and the atrioventricular node, which arise from the embryonic myocardium of the primitive tube as well, maintain, in contrast to the chambers, the above-mentioned embryonic characteristics. Knowledge of the transcriptional gene programs regulating the diversification of the nodal and working myocardial phenotype during development is still incomplete. Research aiming to solve this puzzle not only is driven by pure scientific curiosity regarding the molecular processes underlying the development of the 4-chambered heart of higher vertebrates. Equally important is the insight this fundamental knowledge will give us in the underlying causes of congenital heart defects, which eventually will provide cues for new diagnostic and therapeutic interventions. The most challenging prospect would be to use this knowledge to experimentally differentiate (stem) cells into a specific - nodal or working myocardial - direction in vitro or in vivo. This would give us the tools to generate or renew any cardiac tissue. The work presented in this thesis focuses on the deciphering of the transcriptional pathways regulating the chamber- versus the atrioventricular node- specific gene program.

Chapter 1 starts with a brief introduction for those not familiar with the idiom of the molecular biologist. We discuss several transgenic mouse models, and explain how transcription factors regulate the activity of target genes. Then we review novel insights into the morphogenetic processes and transcriptional gene networks underlying cardiac development, and elaborate on the implications of this new knowledge for our understanding of congenital heart defects. In Chapter 2 we describe a study on the regulatory DNA sequences driving expression of Nppa, the gene encoding atrial natriuretic factor (ANF), during development and disease. Before birth Nppa is expressed specifically in the evolving atrial and ventricular chambers, marking their development. After birth the gene is silenced in the ventricles, where it is reactivated in case of cardiac hypertrophy and heart failure, a feature which makes it an important cardiac biomarker. As a consequence, insight into the regulatory DNA sequences driving Nppa expression, and the transcriptional pathways converging on them, will reveal both the molecular pathways driving chamber development, and those
driving gene expression in the ventricles during cardiac stress. In Chapter 3 we report on our findings on the interplay of three members of the T-box family of transcription factors, Tbx2, Tbx3 and Tbx20, and Bmp/Smad-signaling. In recent years these factors have been shown to play pivotal roles in correct development of the chambers and the atrioventricular canal (the embryonic connection between the atria and the ventricles from which both the atrioventricular node and the mesenchymal cushions, primordia of valves and septa, develop). However, the position in the transcriptional hierarchy of the separate regulators, and the way they interact, were still largely unknown. Combining diverse transgenic approaches, we reveal a novel protein-protein interaction playing a key role in this complex transcriptional gene network. Finally, in Chapter 4, we use regulatory DNA sequences of transcription factor Tbx3 to explore the transcriptional pathways underlying the development of the atrioventricular node, and to get insight into which genes are involved in its formation and function.
Can recent insights into cardiac development improve our understanding of congenitally malformed hearts?

Thomas Horsthuis\textsuperscript{1}, Vincent M. Christoffels\textsuperscript{1}, Robert H. Anderson\textsuperscript{2}, and Antoon F.M. Moorman\textsuperscript{1}

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Abstract

Congenital cardiac malformations account for one-quarter of all human congenital abnormalities. They are caused by environmental and genetic factors. Despite increasing efforts in fundamental research, as yet, the morphogenesis of only a limited number of malformations has been elucidated. Over the last decades, new genetic modifications have made it possible to manipulate the mammalian embryo. Evidence provided using these transgenic techniques has, over the past decade, necessitated re-evaluation of several developmental processes, important in the understanding of normal as opposed to abnormal cardiac development. In this review, we discuss current understanding of the patterning of the initial heart tube, new insights into formation of the atrial and ventricular chambers, and novel information on the origin of the cells that are added to the heart after formation of the initial tube. All of these advances modify our appreciation of malformations involving the venous and arterial poles. As we demonstrate, this new information sheds light not only on normal cardiac development, but also explains the structure of several previously controversial lesions seen in malformed human hearts.
Introduction

Congenitally malformed hearts are the commonest lesions seen in humans at birth, occur in nearly 1% of live births, and are found in up to one-tenth of spontaneously aborted fetuses (Hoffman, 1995). Although many investigators during the previous century speculated on the pathogenesis of these malformations, the precise developmental processes underlying the lesions remained poorly understood. It has been established, nonetheless, that both the

![Diagram of transgenic Cre-loxP system to follow the fate of cells in mice](image)

**Figure 1.** Transgenic Cre-loxP system to follow the fate of cells *in vivo*. (A) Irreversibly to label cells that do or once did express a gene of interest, two transgenic mouse lines are required. In one of the transgenes (mouse I) the coding sequence of Cre (Causes recombination enzyme) is cloned into the locus of the gene of interest. Another transgenic mouse (mouse II) holds a construct in which a ubiquitous promoter (that is, is active in all body cells) potentially drives expression of a reporter gene (here LacZ), however expression does not take place because of the insertion of a stop sequence upstream of the reporter gene. This stop sequence is flanked by loxP sites. When both mice are crossed, in all cells where Cre is expressed, the enzyme recombines the two loxP sites, thus removing the stop sequence and facilitating expression of the reporter gene. As a result, LacZ is active in all cells in which Cre is, or once has been expressed. (See Fig. B and C)
environment (extensively reviewed in Mone et al., 2004) and genetic factors play important causative roles.

With the huge expansion of knowledge and techniques in molecular biology accrued over the last decades of the 20th century, many genetic tools became available which provided important new knowledge. These advances in technology made it possible to generate transgenic mice, revealing the genetic networks underlying both normal and abnormal cardiac development. Using these techniques, it became possible to label irreversibly cells that once expressed a specific gene in the mouse embryo, thus permitting construction of “fate maps” to show the lineage of the marked cells and their progeny (for the interested reader, details of this novel technique are summarized in Figure 1). For the first time, therefore, it became possible to study the origin of the cells of the different components of the mature mammalian four-chambered heart, crucial information when we seek the cause of cardiovascular malformations. The information provided by these studies of cellular lineage often proved discordant with accepted views of cardiac development. Furthermore, the investigations validated in the mouse heart hypotheses made previously on the basis of labelling studies in chick embryos. Thus, it is now accepted that the initial linear heart tube does not contain all precursor cells of the mature heart, but rather gives rise only to part of the mature left ventricle (reviewed in Buckingham et al., 2005). The cells forming the remainder of the mature heart are added subsequently, by additional temporal migrations from the heart-forming areas. Additionally, the studies of lineage showed that the borders between the initial building blocks were not fixed. Rather, the cells initially present in the primitive heart tube differentiate continuously into the more mature myocardium of the cardiac chambers. This means that a structure seen at one particular stage of development will not necessarily retain its cellular composition at subsequent developmental stages. As a result, a malformation found in a specific part of the mature heart might well result from perturbed development of a different early component, or could even be due to abnormal development of components formed outside the embryonic heart.

Many decisions are needed before embryonic cardiac cells develop appropriately from their primitive to their mature phenotype. Many of these decisions are taken, or enforced, by transcription factors. These factors are proteins that are able to bind to specific sequences of DNA (Figure 2). When they bind, together with other transcription factors and co-factors, they turn on, or turn off, the transcription of downstream target genes. These genes, in turn, induce the changes in cellular differentiation. When transcription factors fail to act correctly, cellular differentiation, and hence morphogenesis, are at risk. We now know that many congenital cardiac malformations are caused by mutations that change the protein level or the function of a transcription factor. Since most transcription factors have a role in different processes, act in different parts in the embryo, and at different times of development, such mutations often lead to defects in multiple tissues. In other words, the mutations are pleiotropic.

Taking advantage of this new evidence, we begin this review with a discussion of recent knowledge relative to patterning of the early straight embryonic heart tube. We then
discuss the formation of the chambers by ballooning from this embryonic tube, and the significance of the addition of cells after initial formation of the linear heart tube. We conclude by discussing how these new insights improved our understanding of congenital cardiac malformations, in particular those involving the venous and arterial poles.

The activity of a gene is regulated by activating and repressing transcription factors binding to its promoter.

Figure 2. Schematic presentation of the regulation of the activity of genes by transcription factors. The promoter of a gene is defined as all non-coding DNA sequences which are important for the regulation of its activity, that is, which regulate its transcription into RNA. Transcription factors are DNA-binding proteins that regulate the activity of a gene by binding to specific binding sites in its promoter. The activity of a gene depends on a balance between activating and repressing transcription factors. Transcription factors never act alone, but always with other factors in transcription complexes. Black arrows depict DNA polymerase activity transcribing a gene; green shapes activating transcription factors, red shapes repressing transcription factors.

Patternning of the heart tube

Myocardial cells of the primary heart tube share important physiological characteristics

The heart is the first functional organ that is formed during development. Within 3 weeks of conception in humans, at a stage compatible with the mouse heart on embryonic day 8.5, it is already possible to see the beating heart. The heart needs to be functional from the onset of its formation because the subsequent development of the embryo depends on the circulatory system for the distribution and exchange of nutrients, oxygen, and waste products. At this early stage, the heart is a tube-like structure (Figure 3), which wall consists of an inner endocardial layer, a middle layer of extracellular matrix called the cardiac jelly, and an outer myocardial layer. The myocardium contracts in a peristaltoid fashion (Patten and Kramer, 1933), pumping the blood from the venous to the arterial pole. Already in the 1940s it was recognized that it is the cell-free cardiac jelly which plays an essential role in the pumping function of the tubular embryonic heart (Barry, 1948). All three layers have a mesodermal origin.
All myocytes, being embryonic or adult, share a number of important physiological characteristics (Figure 4 - reviewed in Moorman and Christoffels, 2003). In contrast to adult working myocardial myocytes, the cardiomyocytes of the embryonic heart tube have few contractile elements and a poorly developed sarcoplasmatic reticulum, they display high automaticity and have a low density of gap-junctions; consequently they are poorly coupled.

**Figure 3.** Ballooning model of chamber formation. At embryonic day ED8.0 in mouse, the heart tube consists of primary myocardium sharing important physiological characteristics (see Figure 4). At ED8.5 working myocardium of the ventricular, and only slightly later the atrial chambers, differentiates and balloons out only at the outer curvature of the looping heart. The inflow tract, atrioventricular canal and outflow tract maintain their embryonic phenotype and throughout development are connected at the inner curvature, which also maintains its primary phenotype. The boundary between grey and yellow reflects the pericardial deflection. a, atrium; ea, embryonic atrium; rv, right ventricle; lv, left ventricle; oft, outflow tract; avc, atrioventricular canal; ift, inflow tract; ev, embryonic ventricle.

<table>
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<th>Table: Phenotype of myocardial cells</th>
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<tr>
<td><strong>phenotype</strong></td>
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**Figure 4.** Table with the characteristic properties of the embryonic primary myocardium, the myocardium of the mature nodes, and chamber myocardium. The primary myocardium shares important characteristics with the mature sinus node (SAN) and atrioventricular node (AVN). Both display high automaticity because of high expression levels of the pacemaker channel gene HCN4. Chamber myocardium has a high conduction velocity due to the expression of high levels of fast conducting gap junctions Connexin40 and Connexin43. SR, sarcoplasmic reticulum.
resulting in a slow conduction of the activation wave throughout the heart. Therefore, we
describe the myocardium of the embryonic tube as being primitive, or primary (Figure 4).
Significantly, a number of transcription factors are expressed along the different axes of the
tube, permitting recognition of the patterning of the tube along its cranio-caudal, dorso-
ventral, and left-right axes (Figure 5). The patterning permits phenotypic differentiation of the
different regions of the tube, ensuring its proper function.

**Cranio-caudal patterning**

Patterning along the cranio-caudal axis of the tube (Figure 5) can be recognized from the
distribution of retinoic acid. This patterning is essential for the correct placement of the
ventricles and the atrial chambers formed later in development (Yutzey et al., 1994). The
importance of retinoic acid was shown by experiments in chicken. An excess of retinoic acid
resulted in expansion of developing atrial structures, normally located caudally, while
deficiency led to expansion of the ventricular compartment, which in the normal situation is
positioned cranially (Yutzey et al., 1994; Hochgreb et al., 2003). Equally importantly, there is
electric polarity along the cranio-caudal axis, with the cells showing the highest automaticity
always found at the inflow to the developing heart. This feature insures that the dominant
pacemaker is at the venous pole long before there is formation of a histological discrete
conduction system. The overall effect is to produce slow peristaltoid contraction, which
serves to propel the blood from the inflow to the outflow tract of the heart.

Equally important in determining cardiac design are the different members of the T-
box family of transcription factors (Yutzey et al., 1994; Stennard and Harvey, 2005). One of
the members of the family, *Tbx5*, acts downstream of retinoic acid and shows a similar caudo-
cranial patterning, with the highest expression found caudally (Bruneau et al., 1999). Targeted
knockout of the factor in genetically modified mice reveals its function (for the interested reader, various transgenic possibilities in knock out mice are detailed in our Figure 6). The homozygous mutants show severe malformations, with particularly hypoplasia of the caudal structures of the heart (Bruneau et al., 2001). In humans, the majority of reported TBX5 mutations leads to loss-of-function, and consequently haploinsufficiency of the gene, producing Holt-Oram syndrome, a rare autosomal dominant disorder that is characterized by malformations of the limbs and the heart (Basson et al., 1997). More recently, patients have been observed with a mutation leading to gain-of-function. These subjects show a mild, atypical, phenotype of the Holt-Oram syndrome, but with associated paroxysmal atrial fibrillation (Postma et al., 2008). The cardiac malformations most commonly include defects of atrial and ventricular septation, but many other cardiac malformations have been reported. Among these, abnormalities of the morphologically left ventricle predominate, varying from aberrant trabecular patterning and mitral valvar prolapse to hypoplastic left heart syndrome (Basson et al., 1999). This is in keeping with the caudo-cranial patterning noted in the knockout mice, since as we will discuss, the apical part of the left ventricle balloons from the caudal part of the primary heart tube when considered relative to the right ventricle.

**Figure 6.** Schematic presentation of transgenic mouse models. In the normal situation one gene consists of two active alleles (one from both parents). In a homozygous knockout mouse both alleles have been disrupted and no protein is generated. In a heterozygous knockout mouse one of the alleles has been inactivated. Most often, this causes haploinsufficiency of the gene, i.e. the protein levels of the gene are half of the level in the normal situation. However, in the heterozygous knock out situation, the intact allele might become expressed more actively, resulting in higher than 50% of normal protein levels. Furthermore, if because of polymorphisms the allele that has been knocked out was the most active allele, this might lead to less than 50% of protein levels. Since many genetic disorders in human are caused by mutations resulting in dysfunction of only one of the alleles, heterozygous knockout mice mimic the genetic background of human diseases most closely. In a knock IN mouse, DNA encoding a transgenic gene is cloned into one of the alleles of the targeted gene. Therefore, the transgenic gene (for instance the reporter gene green fluorescent protein (GFP)) is expressed in the same spatiotemporal pattern as the endogenous gene. For clarity, in the model only two homologous chromosomes (represented by grey bars) with one gene (alleles represented by yellow boxes) are shown.
Furthermore, the diversity of the malformations caused by mutation in this single gene is a clear example of a pleiotropic effect.

**Left/right patterning**

The left/right axis of the developing embryo (Figure 5) is established well before formation of the heart tube. Left-right patterning specifies the laterality of most organs within the body, including eventually the heart. Abnormalities in patterning along this axis produce various malformations, including mirror-imagery and the right and left isomeric variants, the latter 2 usually grouped together in the clinical situation as visceral heterotaxy (Capdevila et al., 2000). When there is mirror-imagery, all organs of the body are formed in reverse, in other words with those organs usually formed on the right side retaining their normal anatomy, but being positioned on the left side of the body. In the setting of isomerism, in contrast, the organs on both sides of the body develop with the same morphological characteristics, so that, for example, in right isomerism both lungs have 3 lobes, with each lung supplied by a short bronchus (Figure 7). In the heart, it is only the atrial appendages which can be truly isomeric, since they are the only parts of the atrial chambers dependent on left-right patterning. This is because, as we will explain, both appendages balloon from the same segment of the heart tube, and hence both respond to the same signals. In contrast, the apical parts of the ventricles, these being the components that confer morphological specificity, balloon in series from the original ventral side of the primary tube, with the apical part of the morphologically left ventricle ballooning caudally relative to the apical component of the morphologically right ventricle. Since the ventricles develop in comparable fashion from consecutive parts of the

![Figure 7](https://example.com/figure7.png)

**Figure 7.** The arrangement of most lateralized organs: the lungs and bronchuses, the atrial appendages, liver, spleen, and gut, shown in right and left isomerism. bilat. triang. atrial app., bilaterally triangular atrial appendages; bilat. tubul. atrial app., bilaterally tubular atrial appendages. 1-3, lobes of the lungs.
tube, in molecular terms both have a right and a left side. Hence, both sides of each ventricle respond in comparable fashion to the genes determining laterality.

In molecular terms, evidence of left-right patterning can already be detected during gastrulation, when nodal, a member of the TGFβ family, is expressed only in the lateral plate mesoderm on the left side of the embryo. Its fundamental role in establishment of the bodily axis was shown in transgenic mice expressing nodal on the right side, instead of left-sided mesoderm. These mice developed with complete mirror-imagery, or so-called situs inversus (reviewed in Ramsdell, 2005). The essential regulator of cardiac left-right patterning is Pitx2c (Kitamura et al., 1999; Capdevila et al., 2000), a member of the homeobox gene transcription factor family that acts downstream of nodal. This gene is first detected prior to formation of the heart tube, when it is expressed exclusively in the left heart-forming field. When the heart tube is formed, it is found exclusively in the left part of the looping heart. Mice deficient for Pitx2 have right isomerism of the atrial appendages, common atrioventricular junctions, anomalous pulmonary venous connections, and bilateral systemic venous tributaries (Liu et al., 2002). The mice also show isomerism of the sinus node (Mommersteeg et al., 2007), as do humans with right isomerism (Smith et al., 2006). In humans, mutations in the PITX2 gene are also linked to the Rieger syndrome (reviewed in Amendt et al., 2000), an autosomal dominant disorder that is characterized by abnormalities of the eyes, teeth and umbilicus, but surprisingly not with deficient laterality. This can be explained by the effect of differential gene dosage, with one normal copy of the gene being sufficient to provide appropriate left-right patterning, but insufficient for adequate formation of the teeth and eyes (Liu et al., 2001). Other research with mice has shown that Pitx2 is also expressed in the mesenchymal cells that are added to the heart after formation of the initial heart tube, where it again imposes laterality (Ai et al., 2006; Galli et al., 2008). This finding helps explain the abnormalities of the outflow tract and right ventricle seen in Pitx2-deficient mice, notably double outlet right ventricle or discordant ventriculo-arterial connections, since these structures are almost completely derived from cells added to the heart later in development.

**Dorso-ventral patterning**

When the heart starts to loop, it develops an outer and inner curvature (Figure 3). While cells are continuously added at both sides of the tube, first the ventricles develop at the cranio-ventral side, and only slightly later the atria form more caudally at both dorso-lateral sides. It is the apical parts of the ventricles, and the atrial appendages, which confer morphologic specificity on the ventricular and atrial chambers, respectively. Thus, formation of morphologically specific chambers is regulated not only by cranio-caudal, but also by dorso-ventral patterning (Figure 3 and 5). The principal signal inducing the dorso-ventral axis, however, remains unknown. Based on its pattern of expression, the helix-loop-helix transcription factor Hand1 is a potential candidate (Christoffels et al., 2000). This gene is expressed at the ventral side of the heart tube, and subsequent to looping, along the outer curvature. It is from this side of the loop that the apical parts of the definitive ventricles
balloon, as demonstrated by the expression of genes that are specific for working-myocardium, such as atrial natriuretic factor (anf) and the gap junction protein Connexin40 (Cx40) (Christoffels et al., 2000; Moorman and Christoffels, 2003). When Hand1 is knocked out in mice, the embryos die early in development because of defects of the placental trophoblast (Riley et al., 1998). If the gene is knocked out only in the embryo, and not in the developing placenta, the embryos live until mid-gestation. Cardiac looping does not occur, with the ventricles staying thin-walled, and failing to expand. The chamber-specific gene anf, however, continues to be expressed locally (Riley et al., 1998), indicating that Hand1 is essential for development of the ventricles, but that it is not the primary signal inducing differentiation. Mutations of HAND1 have yet to be detected in humans, most likely because heterozygous deficiency of the gene does not provoke a phenotype, just as has been shown in mice (Riley et al., 1998). An alternative explanation, of course, might be that such mutations are lethal because of extraembryonic defects.
Localized differentiation of working myocardium

Formation of the ventricular and atrial chambers

After its initial formation, the heart tube elongates by the continuous addition of cells at both poles of the heart. These cells are recruited from a highly proliferative pool of mesodermal cells located in the dorsal wall of the pericardial cavity and the pharyngeal arches (Kelly et al., 2001; Cai et al., 2003). Before they differentiate into myocytes, they cease to proliferate (Soufan et al., 2006). The myocytes forming the primary heart tube are in a low proliferative state (Figure 4). Then, only at very localized spots in the tube, the myocytes increase in size and re-initiate cell division (Soufan et al., 2006). Under the control of a chamber-specific genetic programme, these cells develop the properties of chamber myocardium, whilst the remainder of the tube maintains its embryonic, or primary, phenotype (Figure 3 and 4 - Moorman et al., 2000; Moorman and Christoffels, 2003). At the cranial part of the tube, at the original ventral side, primary myocardium begins to differentiate into the working myocardium of the embryonic left ventricle (Figure 3). More caudally, there is differentiation of primary myocardium into the chamber myocardium of the left and right atrial appendages on both dorso-lateral sides (Figure 3). Concomitant with chamber formation, typical apical trabeculated aspects of the chambers develop, much more pronounced in the ventricles than in the atria. The left and right ventricle develop a specific pattern of trabeculations, a feature used by the cardiac pathologist to identify their morphologic identity. The molecular background of this difference between both ventricles is unknown. As the myocytes become converted to a working phenotype, their velocity of conduction increases as they also develop gap-junctions containing Connexin40 and Connexin43 proteins. They also increase their contractile function by building up well-developed sarcomeric and sarcoplasmic reticular structures. These different morphogenetic processes - cell growth, proliferation and differentiation - cause the apical parts of the ventricles, and the atrial appendages, to expand locally. This process of differentiation of the myocardium of the primary heart tube along its outer curvature into the working myocardium of the chambers was initially illustrated by Davis following his studies of the human embryos held in the archive of the Carnegie Institution (Davis, 1927). In the 70ies the expansion from the outer curvature of the looping heart was also illustrated in the textbook of Goor and Lillehei (1975), and furthermore in the textbook of cardiac anatomy produced by Anderson and Becker (1978), albeit using an inappropriate segmental model for the initial heart tube. The concept was then brought to prominence as the ballooning model for formation of the cardiac chambers (Figure 3 - Moorman et al., 2000; Christoffels et al., 2000; Moorman and Christoffels, 2003).

The ballooning model has helped solve one of the major problems confronting generations of cardiac embryologists. This has been to explain how an initially solitary tube, with laminar flow through a single lumen, can transform into a 4-chambered heart in which the pulmonary and systemic circulations work in parallel (Kirby, 2001). The solution to this problem was difficult to understand when it was presumed that the primordiums for all
Figure 8. The segmental model of heart development, in which it was inaccurately assumed that already in the early heart tube all progenitors of the mature building blocks of the heart were present as a linear array of transverse precursor components. sv, sinus venosus. See legend to Figure 3 for other abbreviations.

cardiac chambers were already present in a linear array of transverse segments within the initial heart tube, as in the concept advanced by Anderson and Becker in their atlas of 1978. In this segmental model the atrioventricular canal and the outflow tract were separated by cells fated to become the left and right ventricle, respectively (Figure 8). The major difficulty to overcome was to understand how the mature atrioventricular canal could become connected to the right ventricle, and the mature left ventricle to the subaortic component of the outflow tract. Ingenious but complex solutions had been proposed previously, such as that put forward by Odgers (1938), but the accounts were remarkably difficult to understand. The essence of the ballooning model is the insight that the inflow tract, the atrioventricular canal, and the outflow tract maintain their primary myocardial phenotype, and are interconnected one to another at the inner curvature throughout development. Thus, from the outset the flows through the developing chambers are connected within the primary tube, the parietal wall of the right atrium already being in continuity via the right side of the atrioventricular canal with

Figure 9. Already prior to and during septation, in the looping heart a left and right circulation act in parallel. Electron microscopic picture (A), model without cushions (B) and with atrioventricular and outflow tract cushions (C) of a frontal view of a ED10 septating mouse heart. In the models the outflow tract has been shifted sideward for clarity. The dotted arrows indicate the left (systemic) and right (pulmonary) circulation acting in parallel. In the models the cardiac cushions are indicated in yellow, cush, cardiac cushion. See legend to Figure 3 for other abbreviations.
the wall of the developing right ventricle, and the wall of the presumptive subaortic outlet already being in continuity, via the inner heart curvature, with the roof of the developing left ventricle. Long before overt cardiac septation, therefore, the systemic and pulmonary circulations are properly arranged in parallel (Figure 9 shows this parallel arrangement in the septating heart).

**Chamber formation requires a network of activating and repressing transcription factors**

We are just beginning to understand the molecular mechanisms underlying the strictly localized activation of the chamber-specific genetic programme. By an interplay of transcription factors, including Nkx2-5, Gata4, and the T-box transcription factors Tbx5 and Tbx20, the chamber-specific programme is potentially activated throughout the entire heart tube (Figure 10), inducing differentiation of primary into working myocardium (Bruneau et al., 2001; Hiroi et al., 2001). Formation of chamber myocardium, nonetheless, is prevented in the inflow tract, atrioventricular canal, inner curvature and outflow tract by the transcription factors Tbx2 and Tbx3, which repress the chamber-specific genetic programme (Figure 10 and 11 - Habets et al., 2002). Tbx2 is expressed from early stages onwards when formation of the chambers commences, and decreases prior to birth. Tbx3, however, is expressed until adulthood, and delineates the developing central conduction system (Hoogaars et al., 2004).

This mechanism of localized complementary repression has important consequences on our thinking about congenital malformations that might occur during formation of the chambers. If repression is incomplete, chamber-specific genes will be expressed ectopically,

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**Figure 10.** Model for the mechanism of transcriptional activation and repression of chamber genes. Broadly expressed transcription factors like GATA, Nkx2-5, Tbx5 and Tbx20 potentially activate chamber-specific genes throughout the looping heart. Tbx2 and Tbx3 are specifically expressed only in the forming inflow tract, atrioventricular canal, inner curvature and outflow tract. These transcriptional repressors compete with Tbx5 for DNA binding and for Nkx2-5 and as a result prevent the activation of the chamber-specific gene program in these regions. Black oval depicts DNA polymerase. See legend to Figure 3 for abbreviations.
resulting in a chamber-like phenotype in the developing inflow tract, atrioventricular canal, inner curvature and outflow tract. Because, as we will explain later, some of these structures give rise to the mature nodes of the conduction system, ectopic expression of chamber-specific genes in these tissues might also lead to arrhythmias. For instance, changes in the essential low electrical coupling between the sinus node and the adjacent atrial myocardium may cause dysfunction of the sinus node. When rapidly conducting gap junctions, such as those containing Connexin40 or Connexin43, become ectopically expressed in the sinus node, the negative resting membrane potential of the atria (-90 mV) will hyperpolarize the more depolarized membrane potential of the small sinus node (-60 mV). The electrical load from the atrial myocardium on the sinus node will slow the rate of diastolic depolarization, and hence spontaneous activity. Should the electrical load become sufficiently large, it may even lead to sinusatrial arrest. Likewise, increased conduction in the atrioventricular node will reduce the conduction time between the atrial and ventricular chambers. Consequently, all four chambers will contract almost simultaneously, which will impair the normal pump function of the heart. An additional disadvantage is that supraventricular tachycardias will be more easily conducted to the ventricles, thereby potentially causing life-threatening ventricular tachycardias. In addition, when the signaling pathways in the primary myocardium are disturbed, it is possible that this may induce formation of rapidly conducting atrioventricular muscular connections across the malformed atrioventricular junctions. Recent experiments from our laboratory, as yet unpublished, show that these accessory tracks can lead to pre-excitation of the ventricles, as seen in humans with Wolff-Parkinson-White syndrome.

In Tbx2-deficient mice, chamber-specific genes are expressed throughout the developing atrioventricular canal (Harrelson et al., 2004). Although an atrioventricular constriction was visible in these mice, it was less than normal, with compromised formation of the endocardial cushions. When Tbx3 was knocked out, mice died before birth, but initially
an obvious macroscopic cardiac phenotype was not reported (Davenport et al., 2003). More recent investigations have shown that, in Tbx3 knock out mice, the gap junction genes Connexin40 and Connexin43 are ectopically expressed in the sinus node (Hoogaars et al., 2007), whereas the bundle branches fail fully to develop (Bakker et al., 2008). Moreover, these mice exhibit ventricular septal defects and double outlet right ventricle. Functional consequences of the genes expressed ectopically in the His bundle and bundle branches, which are normally exclusively expressed in the working myocardium, surprisingly could not be detected (Bakker et al., 2008). The functional consequences for the sinus node, in other words whether these mice suffer from bradycardias or sinuatrial arrests, have yet to be investigated. In humans, mutations of TBX3 cause the ulnar-mammary syndrome (Bamshad et al., 1997). This syndrome is characterized by defects of the upper limbs on the ulnar side, malformations of the external genital organs, hypoplasia of the mammary glands, and malformations of the teeth (Bamshad et al., 1999). Only very rarely have malformations of the heart been reported, such as ventricular septal defect at birth, and pulmonary stenosis at a later age (Meneghini et al., 2006). Again unexpectedly, thus far no instance has been encountered of abnormal conduction. This relatively mild cardiac phenotype cannot be explained by the redundant expression of Tbx2, because Tbx2 is neither expressed in the sinus node, nor in the atroventricular bundle. Apparently, half of the normal levels of the TBX3 protein in the embryo are sufficient for appropriate development of the heart, but not the limbs. To the best of our knowledge, mutations of TBX2 have not yet been reported in man.

Further formation and septation of the atrial chambers

We have explained how ballooning of the appendages from the original caudo-dorsal part of the primary heart tube provides the basis for formation of the right and left atrial appendages. We have also explained that, at this stage, the remainder of the atrial component of the heart tube retains its primary myocardial phenotype (Figure 4). At this stage of development, the systemic venous tributaries drain to the caudal pole of the tube in symmetrical fashion (Figure 12A and B), and as yet there is no formation of either the lungs, or the pulmonary vein. The caudal part of the tube, nonetheless, remains attached to the pharyngeal mesoderm via the dorsal mesocardium. It is through the dorsal mesocardium that new material is added to the venous pole of the developing heart. This newly added mediastinal myocardium functionally is working myocardium because it expresses fast conducting channels composed of Connexin40. However, it is different from the working myocardium of the appendages because it does not express atrial natriuretic factor (Figure 12). In the mouse, the mediastinal myocardium is initially restricted to the area around the persisting dorsal mesocardium (Figure 12 - Soufan et al., 2004). With ongoing development, the area formed by mediastinal myocardium expands considerably, and comes to form the larger part of the body of the developing left atrium, the primary atrial septum, and a significant part of the developing right atrium up to the left venous valve. At the same time, there is a marked realignment of the systemic venous tributaries, such that the left sinus horn diminishes in size, with the left
cranial cardinal vein becoming incorporated, in the mouse, into the left atrioventricular groove, draining to the right atrium as well (Figure 12C and D). This then sets the scene for atrial septation, since the developing pulmonary vein opens to the left atrium through a solitary opening in the dorsal mesocardium, between the so-called pulmonary ridges, adjacent
to the developing atrioventricular junction. The primary atrial septum then grows from the newly formed mediastinal myocardial component, approaching the endocardial cushions which themselves are dividing the atrioventricular canal. As the primary atrial septum grows towards the cushions, it carries on its leading edge a mesenchymal cap (Figure 12). It is then fusion of the mesenchymal cap with the fused atrial aspect of the endocardial cushions that closes the primary atrial foramen. Prior to closure of the primary atrial foramen, however, the upper margin of the primary septum breaks down so as to permit the richly oxygenated placental blood returning to the heart through the umbilical and eventually the caudal cardinal vein to continue to reach the left side of the developing heart.

The point of fusion of the mesenchymal cap and the endocardial cushions is then further reinforced by growth of non-endocardium derived mediastinal mesenchyme through the mesocardium into the heart, the so-called vestibular spine or dorsal mesenchymal protrusion (His, 1880; Webb et al., 1998; Mommersteeg et al., 2006; Snarr et al., 2007). We now know that, subsequently, this non-endocardially derived mesenchyme is muscularised to form the basal buttress of the atrial septum (Mommersteeg et al., 2006). The so-called “septum secundum”, however, is no more than a fold in the atrial roof. Indeed, in the human heart this superior interatrial fold is not developed until after the solitary pulmonary vein has migrated to the roof of the left atrium, achieving separate opening for the four pulmonary veins in the process (Webb et al., 2001). The “septum secundum” is then simply the deep fold between the connections of the caval veins to the right atrium, and the pulmonary veins to the left atrium. This concept of atrial septal development also shows why sinus venosus and coronary sinus defects are interatrial communications, rather than atrial septal defects, since they are outside the confines of the atrial septum, this being limited to the floor of the oval fossa and the basal buttress formed by muscularisation of the vestibular spine (Anderson et al., 2004).

Primary myocardium signals to adjacent endocardium to form cardiac cushions and later in development gives rise to the mature nodes of the conduction system

Subsequent to ballooning of the atrial appendages and the apical parts of the ventricles, it becomes possible to recognize the atrioventricular canal as the component of the primary tube between the developing atrial and ventricular components. Because of the differences in growth, the inflow tract, atrioventricular canal, and the outflow tract are the persisting components of the primary tube roofed by the primary myocardium of the inner curvature. Responding to signals from this myocardium, the endocardial cells within the canal undergo a process of epithelial-to-mesenchymal transformation, and form the atrioventricular endocardial cushions (Figure 9 - Eisenberg and Markwald, 1995). Later in development, these cushions give rise to some of the leaflets of the mitral and tricuspid valves (de Lange et al., 2004). Furthermore, during development cushion tissue fuses at the atrioventricular border with epicardial tissue of the atrioventricular groove, insulating the atria from the ventricles.
electrically (Wessels et al., 1996). In the adult heart the fibrous body only consists of cells from the atrioventricular cushions (de Lange et al., 2004) and forms an integral part of the atrioventricular septum (Kanani et al., 2005). Additional cushions are formed within the outflow tract, which act initially as septal structures (Kanani et al., 2005), but eventually cavitate along their distal margins to form the leaflets of the arterial valves and their supporting sinuses (de Lange et al., 2004). The proximal parts of the cushions, subsequent to their fusion, muscularise to form the subpulmonary infundibulum, which eventually is converted into a free-standing muscular sleeve. The outflow cushions, therefore, play a crucial role in separating the systemic and pulmonary circulations, albeit that no septal structures remain in the definitive heart subsequent to closure of the embryonic interventricular foramen (Anderson et al., 2003).

Concomitant with formation of the chamber myocardium, it is possible to detect a mature electrocardiogram (Figure 13). As we have already discussed, the impulse is generated by the pacemaker cells at the venous pole, which then induces a rapid depolarization of the atrial myocardium. Subsequently, there is a delay in propagation of the impulse through the slowly conducting primary myocardium of the atrioventricular canal, followed by rapid propagation through the ventricles. Later in development, parts of the primary myocardium of the inflow tract and atrioventricular canal become the sinus and the atrioventricular nodes of the definitive conduction system, respectively (Figure 13 - Moorman and Christoffels, 2003). These mature nodes share essential phenotypic characteristics with their embryonic primordiums (Figure 4). The remaining parts of the inflow corridor, the atrioventricular canal, and the outflow tract do not contribute to the histologically discrete conduction system, but instead differentiate into the working myocardium of their appropriate chambers (Rana et al., 2007), albeit that their primary origin can offer explanation for several known cardiac arrhythmias (Moorman et al., 2005).

![Figure 13](image-url)

**Figure 13.** The embryonic myocardium of the inflow tract and atrioventricular canal give rise to the mature sinus node and atrioventricular node, respectively. (A) ED9.5 mouse embryo, with an ECG recording drawn in it. As soon as the chambers form, an “adult-like” ECG can be detected (See text for details). Blue line marks the contour of the atria, red line the ventricles. Arrow gives direction of the action potential. (A,B,C) The inflow tract (dark green dot or circle in A,B) gives rise to the mature sinus node (dark green circle in C), whereas the atrioventricular canal (light green dot or circle in A,B) gives rise to the mature atrioventricular node (light green circle in C). See legend to Figure 3 for abbreviations.
**Subsequent to formation of the linear heart tube, cells are continuously added to the heart**

Already from 1969 onwards, various labelling studies in chicken suggested that cells were added to both poles of the growing heart from the surrounding mesoderm (Figure 14 - Stalsberg and de Haan, 1969; Virágh and Challice, 1973; De la Cruz et al., 1989; De la Cruz and Sanchez-Gomez, 1998b). Since at that time cardiac development tended to be explained on the basis of the segmental model (Figure 8), in which the primordiums of all future components were postulated to be present in the initial heart tube (Rosenquist, 1970; Garcia-Martinez and Schoenwolf, 1993), these important findings were largely ignored. In 2001, however, studies in chicken (Waldo et al., 2001; Mjaatvedt et al., 2001), and use of transgenic fate mapping techniques in mouse (Kelly et al., 2001), showed conclusively that the cells forming most of the mature right ventricle and outflow tract were added to the heart subsequent to the initial stage of looping. It was then shown by use of other genetically modified mice that cells were also added at the venous pole (Cai et al., 2003; Christoffels et al., 2006). It is now generally accepted that the progenitors within the initial linear tube give rise only to most of the apical part of the mature left ventricle (reviewed in De la Cruz and Markwald, 1998a; Zaffran et al., 2004). In the earliest stage, cells can be added via the mesocardium, which connects the linear tube with the dorsal pericardial wall over its full length. During looping, the mesocardium breaks to form the transverse sinus of the

**Figure 14.** Subsequent to formation of the initial heart tube, extracardiac cells are added to the heart continuously. Left lateral side of ED9.5 mouse embryo with contours of the heart drawn in it. Red arrows depict mesodermal cells from the second heart field which are added to the venous and arterial poles by recruitment. Yellow arrows depict cardiac neural crest cells which are added by migration after formation of the initial heart tube. Blue arrow depicts precursor cells of the sinus venosus. See legend to Figure 3 for abbreviations.
pericardium. Subsequent to this breakdown, the tube is connected to the body wall only at the
venous and arterial poles. This enables the heart to beat freely within the pericardial cavity. After
the breakdown of the mesocardium, nonetheless, cells can then only be added at the
venous or arterial poles of the growing heart. We now know that it is through these poles that
further additions are made to the developing heart.

**The Second Heart Field**

Progenitor cells derived from splanchnic mesoderm bordering the initial heart tube
continuously differentiate to myocardium, in a process nowadays often referred to as
recruitment (red arrows in Figure 14). This causes the heart tube to elongate at both sides.
After the initial differentiation into embryonic myocardium, cells will undergo additional
differentiation, thus contributing to different building blocks of the heart. During recruitment,
the cells do not move individually or as clusters, but are taken up as sheets, without changing
position relative to their neighbours (Stalsberg and de Haan, 1969). The cells are recruited to
the initial linear tube from a region in the dorsal wall of the pericardial cavity that has now
become known as the second heart field (reviewed in Buckingham et al., 2005). In mouse, the
cells from this field are characterized by their expression of the LIM homeodomain
transcription factor *Islet1* (Cai et al., 2003). The so-called second heart field is used in this
fashion to distinguish the cells of the initial linear tube, which are considered to be derived
from a primary heart field. We are comfortable with this nomenclature, although along with
others (Abu-Issa et al., 2004), we question the discreteness of the alleged fields (Moorman et
al., 2007). This scepticism is, perhaps, reinforced by the recent finding that, in early stages of
development, the Islet1 protein can also be detected in the progenitors of the primary heart
field among species (Yuan and Schoenwolf, 2000; Prall et al., 2007; Brade et al., 2007).
These findings render Islet1 a pan-cardiac marker, and now leave the purported second field
without any specific marker, supporting the notion of the existence of a single heart field,
from which precursor cells are gradually added to the heart in a temporal sequence. It has
been shown, nonetheless, that cells from the cranial part of the purported second field express
the fibroblast growth factor gene Fgf10, and it is this population which gives rise to the
outflow tract and the right ventricle (Waldo et al., 2001; Mjaatvedt et al., 2001; Kelly et al.,
2001). In contrast, most of the atrial myocardium is derived from the more caudal part of the
field (Figure 12 and 14 - Cai et al., 2003; Galli et al., 2008). It remains a fact that only the left
ventricle is derived from progenitors of the first heart field, which give rise to the initial linear
heart tube early in development.

The identification of the second heart field, and its derivatives, has obvious
implications for our thinking about congenital cardiac defects. First, it shows that the left and
right ventricles have a distinctive transcriptional history. A biopsy of the left ventricle,
therefore, does not necessarily tell us anything about the right ventricle! It may also explain
why cardiac defects are localized to one side of the heart, albeit that disordered flow
subsequent to the completion of septation provides an equally convincing explanation of
lesions such as hypoplastic left heart syndrome or pulmonary atresia with intact ventricular septum. The migration of temporally distinct populations of cells from the heart-forming areas, nonetheless, does show that the primary cause of malformations of the outflow region and right ventricle, varying from functionally univentricular hearts with dominant left ventricles to subtle defects of the outflow tracts, might be located outside the linear heart tube, and be located in the so-called second heart field. DiGeorge syndrome, for example, is a dominant disorder that is linked to a micro-deletion on human chromosome 22q11. It is characterized by a spectrum of abnormalities involving the outflow tracts, such as common arterial trunk or tetralogy of Fallot, and extra-cardiac defects like craniofacial dysmorphologies and hypoplasia of the thymus and parathyroid glands. This spectrum of abnormalities can be attributed to abnormal development of the pharyngeal arches and second heart field (Scambler, 2000). Because the T-box factor *Tbx1* was shown to be both located within this chromosomal deletion, and expressed in the developing pharyngeal arches and second heart field, it was identified as a candidate gene. Indeed, in 2001 it was shown that mice deficient for *Tbx1* phenocopy important aspects of the DiGeorge syndrome, including the malformations involving the outflow tracts (Jerome and Papaioannou, 2001). Also, since the atrial chambers, and the right ventricle, are largely derived from the second heart field, it has been suggested that mutations of genes that are important for correct regulation of the second heart field may lead to congenital malformations at both poles. Indeed, mice lacking the transcription factor *Islet1* display severe malformations of the outflow tract and right ventricle, but also of the atrial chambers (Cai et al., 2003). In humans, however, defects that involve both the inflow and outflow regions of the heart are very rare. This rarity strongly suggests that a defect within the highly proliferative progenitors of the second heart field, sufficiently powerful to disturb both poles of the heart, is lethal.

**Cells from the neural crest migrate into the arterial pole to aid in separation of the pulmonary and aortic pathways**

Whereas the mesodermal cells are added to the heart by recruitment, cardiac neural crest cells are added to the arterial pole by migration (yellow arrows in Figure 14). After the neural tube and neural crest have been formed from the ectoderm during neurulation, the cells of the neural crest differentiate into ectomesenchyme, breaking up into individual cells that subsequently migrate through the body to give rise to diverse tissues such as neural ganglions, melanocytes, and the adrenal glands. The ectomesenchymal cells of the cardiac neural crest migrate into the outflow tract, populating the outflow cushions formed by endothelial-to-mesenchymal transformation. The cushions, packed by the cells from the neural crest, initially form a septal structure throughout the lumen of the initially solitary outflow tract. They subsequently contribute to the formation of the arterial valvar leaflets and their supporting sinuses, and the most proximal parts muscularise to form the free-standing subpulmonary infundibulum (Anderson et al., 2003). The cells initially separating these structures, however, subsequently disappear, so that eventually there are no septal components between the arterial
valves, or between the subpulmonary infundibulum and the aortic root (Webb et al., 2003). The cardiac neural crest cells also give rise to the connective tissue of the thymus, thyroid gland and parathyroids. Ablation of the cardiac neural crest cells in the chick has been shown to cause malformations such as common arterial trunk, and non-cardiovascular defects like malformations of these glands in the neck (reviewed in Hutson and Kirby, 2003). Indeed, deficiency of the cardiac neural crest mimics the features from DiGeorge syndrome, which has been linked to the \( Tbx1 \) gene. It is nowadays established that, although not the principal perpetrator of the syndrome, disruption of migration of cells from the cardiac neural crest, either directly or indirectly by impairment of the \( Tbx1 \) dose, is an important mediator of the typical phenotype.

**Conclusions**

The ability genetically to modify mice has rapidly increased our insights into the mechanisms underlying cardiogenesis. These techniques have improved our understanding of several developmental mechanisms essential for shaping the normal developing heart. Awareness of these mechanisms, and of the transcriptional networks which regulate them, provides vital new clues for understanding congenital cardiac malformations. Based on these insights, it has been possible to detect novel candidate genes. Genetic and molecular research exploring these genes has shown them to produce congenitally malformed hearts. This new knowledge, however, should also serve to makes us humble, since the transcriptional networks and cellular interactions regulating cardiac development are shown to be remarkably complex. We have only just started to unravel them.

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Distinct regulation of developmental and heart disease-induced ANF expression by two separate distal sequences

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Abstract

*Nppa*, encoding atrial natriuretic factor, is expressed in fetal atrial and ventricular myocardium, and is down-regulated in the ventricles after birth. During hypertrophy and heart failure, *Nppa* expression is re-activated in the ventricles, and serves as a highly conserved marker of heart disease. The *Nppa* promoter has become a frequently used model to study mechanisms of cardiac gene regulation. Nevertheless, the regulatory sequences that provide the correct developmental pattern and ventricular re-activation during cardiac disease remain to be defined. We found that proximal *Nppa* fragments ranging from 250 bp to 16 kbp provide robust reporter gene activity in the atria, and correct repression in the atroventricular canal and the nodes of the conduction system *in vivo*. However, depending on fragment size and site of integration into the genome of mice, the fetal ventricular activity was either absent or present in an incorrect pattern. Furthermore, these fragments did not provide ventricular re-activation in heart disease models. These results indicate that the proximal promoter does not provide a physiologically relevant model for ventricular gene activity. In contrast, two modified bacterial artificial chromosome clones with partially overlapping genomic *Nppa* sequences provided appropriate re-activation of the green fluorescent protein reporter during pressure overload induced hypertrophy and heart failure *in vivo*. However, only one of these bacterial artificial chromosomes provided correct fetal ventricular green fluorescent protein activity. These results show that distinct distal regulatory sequences, and divergent regulatory pathways, control fetal ventricular activity and re-activation of *Nppa* during cardiac disease, respectively.
Introduction

A variety of mechanical, hormonal and genetic stimuli can cause cardiac enlargement and eventually heart failure. In response to these stimuli the ventricular myocytes increase in size and activate a hypertrophy response gene program. This gene program is characterized by induction of cardiac contractile protein genes and re-activation of a ‘fetal’ gene program, normally only active during cardiac development (reviewed in (Olson and Schneider, 2003;Chien et al., 1991)). Of the known genes that are re-activated, Nppa, encoding atrial natriuretic factor (ANF), is probably the best characterized (Chien et al., 1991;Rosenzweig and Seidman, 1991). Nppa is expressed specifically in the myocardium of the atria and ventricles of the embryonic and fetal heart, and is the first marker of their formation (Zeller et al., 1987;Christoffels et al., 2000). After birth, Nppa expression is down-regulated in the ventricles, where it is re-activated again in situations of cardiac stress.

Proximal promoter fragments (0.5-3.4 kbp) of Nppa of human, rat, mouse and Xenopus have been shown to drive atrial and fetal ventricular expression both in vivo (Field, 1988;Seidman et al., 1991;Knowlton et al., 1995;Habets et al., 2002;Small and Krieg, 2003;von Harsdorf et al., 1997) and in cell cultures (Durocher and Nemer, 1998;Seidman et al., 1988), and to provide a hypertrophy stress response in cardiomyocyte cell cultures and after injection as plasmid DNA in the ventricles of failing dog hearts (Ardati and Nemer, 1993;Shubeita et al., 1990;von Harsdorf et al., 1997). As such, the proximal Nppa promoter has become the most widely used model promoter that has importantly contributed to revealing transcriptional networks involved in cardiac gene regulation during cardiac development, health, and disease (reviewed in (Nemer and Nemer, 2001;Temsah and Nemer, 2005;Houweling et al., 2005)). Nonetheless, the Nppa promoter fragments have been shown to lack hypertrophy responsiveness in vivo (Knowlton et al., 1995). From these findings it has been concluded that the fetal gene program is regulated by sequences within the 0.7 kbp proximal promoter fragment, while the re-activation of the 'fetal' gene program during disease is regulated by sequences residing outside this fragment and therefore by a divergent pathway. However, a recent study indicated that the ventricular activity is absent from an even larger promoter fragment (de Lange et al., 2003), hindering any conclusions regarding the divergence of fetal and hypertrophy pathways. In addition, this lacking function would make the proximal promoter a physiologically less relevant read-out for the activity of cardiac transcriptional pathways. To address these issues, we assessed the regulatory activity of the Nppa locus in transgenic mice in detail. We found that the proximal Nppa promoter fragments, currently frequently used in studies of cardiac development and disease, lack critical regulatory functions for ventricular activity during development and in disease. Furthermore, the fetal activity of Nppa is regulated independently from its stress-response and re-activation during cardiac disease, both processes requiring distal regulatory sequences. Therefore, ventricular activity before birth and re-activation during disease are regulated by divergent transcriptional pathways.
Materials and methods

Transgenic mice

The transgenic promoter-reporter lines -0.7rLacZ and -3/+4mCre, and the αMHC-Gal4 heart failure mouse model have been described previously (Habets et al., 2002; de Lange et al., 2003; Habets et al., 2003). The -0.7rLacZ construct was targeted as a single copy to the Hprt locus as described (Hprt-0.7rLacZ) (Figure 1A-B) (Bronson et al., 1996). To generate the -0.7mLacZ construct we replaced the rat Nppa promoter sequence by that of the mouse Nppa promoter sequence in the -0.7rLacZ construct (Figure 1D). Subsequently, to make the -0.7/+4mLacZ construct (Figure 1D), we replaced upstream sequences in the -3/+4mCre fragment with this -0.7mLacZ construct. The -3mCre construct was generated by truncating the -3/+4mCre construct in the third exon, proximal from the NRSE (Figure 1D). A -380/-138 rat Nppa promoter fragment was PCR amplified and fused to the -230/+126 cTnI-LacZ construct flanked by insulator sequences previously described (Habets et al., 2002), to generate the 0.25rNppacTnILacZ construct (Figure 1D).

Two BAC clones of a mouse 129 SvJ BAC library (Incyte, St. Louis, MO) harbouring Nppa were end-sequenced to establish the genomic sequences they contain. BAC clone 337 ranges from -27 kbp to +127 kbp, clone 336 from -141 to +58 kbp relative to the transcription start site of Nppa (Figure 1C-D). At the translation start site of Nppa of both BACs, we replaced sequence cc.cac.gcc.agc.ATG.ggc by Egfp, using the BAC modification protocol kindly provided by Shiaoching Gong and Nathaniel Heinz (Gong et al., 2002). Subsequently, the -11/+5mEgfp construct was generated from the modified BAC336-Egfp construct (Figure 1D).

Supplementary methods are available online and at the end of this chapter.

Results

The activity of the proximal Nppa promoter is context dependent

To assess the spatio-temporal activity profile of the widely used 0.7 kbp Nppa promoter fragment, it was targeted as a single copy to the hypoxanthine phosphoribosyltransferase (Hprt) locus (Figure 1A, B). This locus is transcriptionally accessible throughout development and adult life, allowing promoter constructs to retain their activity and specificity, while variations in activity resulting from variation in copy number and site of insertion in the chromatin are ruled out (Bronson et al., 1996). Expression of the lacZ reporter driven by the promoter was restricted to the heart throughout development (Figure 2). Onset of transgene expression was observed at E8 at the ventral side of the heart tube (not shown), comparable to the onset of Nppa expression (Christoffels et al., 2000). From E9.5 onwards both the reporter gene and the endogenous Nppa gene were expressed in atrial and ventricular
Figure 1. Overview of all Nppa promoter constructs used in this study. (A) Targeting vector used to target the 0.7 kbp rat Nppa promoter to the Hprt locus. (B) Southern blot for verification of single integration at the Hprt locus after BamH1 digestion of genomic DNA. 1, wild type; 2, wild type; 3, hemizygous male of line A; 4, homozygous female of line A; 5, hemizygous male of line B; 6, heterozygous female of line B. The position of the RsaI probe used for Southern blotting is indicated (dotted line). (C) Web based RankVISTA alignment of human against mouse Nppa genomic region covered by BAC336 and BAC337. BAC336 and BAC337 share 85 kbp of overlapping sequences and both contain unique sequences either up- or downstream, respectively. RankVISTA bars depict evolutionary conserved segments where the heights scale with statistical significance [-log10(P-value)]. (D) Constructs used to generate transgenic mice. The table indicates the number of founders that express the transgene (Expression) over the total number of founders or lines obtained (Transgenes). Grey lines in panel A indicate mouse Hprt locus sequences. Thick black lines indicate Nppa promoter sequences, green thick lines the 0.25 kb Nppa ‘atrial’ regulatory module, the thick grey line and box in panel D the -230/+126 cTnI promoter fragment. Boxes represent exons, dark blue boxes the reporter gene and the red box the human growth hormone polyadenylation signal. CNCS, conserved non-coding sequence; UTR, untranslated region.
chamber myocardium, but not in the atrioventricular canal and outflow tract (Figure 2; 3; 7). Nppa expression is absent from the sinoatrial and atrioventricular node (Wharton et al., 1988; Houweling et al., 2002). Analysis of E17.5 embryos revealed that also the Hprt-0.7rLacZ construct was not active in these components (Figure 3J-L). However, the transgene was ectopically active in the sinus horns and mediastinal atrial myocardium (Figure 2G-I; 3I). After E9.5 ventricular Nppa expression becomes restricted to the trabeculated myocardium, and after birth it largely disappears (Figure 3). In contrast, before and after birth the Hprt-0.7rLacZ construct remained active in a transmural pattern in the ventricles (Figure 3; 4A, D). These findings indicate that while the 0.7 kbp promoter drives important aspects of the pattern of Nppa, it lacks regulatory sequences for the correct fetal ventricular pattern and postnatal down-regulation, and for correct repression in the mediastinal atrial myocardium and sinus horns.

The pattern of the Hprt-0.7rLacZ construct was compared to that of two lines in which the -0.7 kbp promoter-lacZ construct was randomly integrated in the genome. Both lines showed similar cardiac expression during development, including ectopic expression in the sinus horns and mediastinal myocardium, albeit that the ventricular activity was relatively weak. In addition, the -0.7rLacZ promoter fragment was ectopically active outside the heart (Figure 2). At birth, line A (-0.7rLacZA) expressed lacZ homogeneously in the atria, whereas some activity was present in the left ventricle (Figure 4B). In the adult heart the atrial expression had become patchy and the ventricular activity had disappeared (Figure 4E). In line B, expression in the atria was patchy at day of birth, while expression in the ventricles was not detectable anymore (Figure 4C). In the adult hearts of this line, only a few atrial cells still expressed the construct (Figure 4F). These observations indicate that the previously observed postnatal ventricular down-regulation of the proximal Nppa promoter fragments (Knowlton et al., 1995) may not be specific to the ventricle, and dependent on the site of integration.

**Key regulatory functions of Nppa reside within a small ‘atrial’ module, whereas extension of proximal promoter fragments diminishes ventricular activity**

Previous studies indicated that the 0.7 kbp Nppa promoter fragment is organized in three modules, a ventricular and developmental module, an atrial module and a basic cardiac promoter (Argentin et al., 1994; Durocher and Nemer, 1998). We tested the contribution of the ventricular module in vivo, by placing a 0.25 kbp fragment containing only the atrial module upstream of a cTnI promoter fragment (0.25rNppacTnI) (Figure 1D) that normally is always expressed in the atrioventricular canal and only shows limited expression in the atria (Habets et al., 2002). This resulted in an expression profile comparable to that of the 0.5NppacTnI fragment containing both the atrial and the ventricular module (Figure 7; Supplementary Figure 1; Habets et al., 2002), with repression of the cTnI promoter fragment in the atrioventricular canal, and activity driven in the atria and in the left ventricle. These findings
indicate that the regulatory sequences responsible for the atrial and ventricular activity of the 0.7 kbp proximal \textit{Nppa} promoter reside within the atrial module, and that the ventricular module does not contribute significantly.

\textbf{Figure 2.} Endogenous \textit{Nppa} expression compared to the targeted \textit{Hprt-0.7rLacZ} and the randomly integrated \textit{-0.7rLacZ} construct. (A, D, G) Whole mount \textit{in situ} hybridisation of \textit{Nppa}. (B-C, E-F, H, I) Whole mount \(\beta\)-galactosidase staining. Both the \textit{Hprt-0.7rLacZ} and the \textit{-0.7rLacZ} constructs showed ectopic expression in the sinus horns (arrow heads in H, I). ra, right atrium; la, left atrium; rv, right ventricle; lv, left ventricle; oft, outflow tract; avc, atrioventricular canal; lsh, left sinus horn; rsh, right sinus horn.

We previously showed that a 7 kbp mouse \textit{Nppa} fragment (-3/+4mCre), that includes the 0.7 kbp fragment, is virtually inactive in the ventricles (Figure 7; Supplementary Figure 2A; de Lange et al., 2003). By truncation experiments we tested whether a ventricular repressor either upstream (located between -3 kbp and -0.7 kbp) or downstream (located between -0.7 and +4 kbp) could explain the lack of ventricular activity. Both the up- and downstream truncated construct (Figure 1D) showed almost exclusively atrial expression, similar to -3/+4mCre transgenes (Figure 7; Supplementary Figure 2A-C). These findings argue against a repressor located within the 7 kbp fragment that inhibits ventricular expression.

The 7 kbp Nppa fragment is from mouse, whereas the 0.7 kbp promoter fragment that showed activity in the ventricles is from rat. The rat proximal promoter contains marked sequence differences with those of mouse and human (Durocher and Nemer, 1998). To examine whether these differences could account for the lack of ventricular activity in the
larger constructs, we tested the 0.7 kbp of the mouse Nppa promoter (-0.7mLacZ) (Figure 1D). Similar to the rat 0.7 kbp promoter, this construct was active in both the atria and the ventricles, was correctly inactive in the atrioventricular canal and outflow tract, and was ectopically active in the sinus horns (Figure 7; Supplementary Figure 2D). Thus, the lack of ventricular expression of the larger constructs cannot be explained by a species difference.

In search for lacking sequences responsible for ventricular activity we subsequently generated a larger construct of 16 kbp (-11/+5mEgfp) (Figure 1D). However, this construct appeared completely inactive in the ventricles as well (Figure 7; Supplementary Figure 2E). Taken together, our data indicate that enhancers outside a -11 kbp to +5 kbp fragment are required for ventricular activity.

Figure 3. Comparison of expression patterns of endogenous Nppa and the Hprt-0.7rLacZ transgene. (A, C, E, G, J) Section in situ hybridisation of Nppa. (B, D, F, H, I, K-L) Section β-galactosidase staining. san, sinoatrial node; avn, atrioventricular node; mm, mediastinal myocardium; rsv, right systemic vein; tr, trabeculated myocardium; c, compact myocardial layer. See legend to Figure 2 for other abbreviations.

Distal regulatory DNA regions for the ventricular pattern and fetal activity

To identify distal regulatory sequences we used two Nppa containing BAC clones of 150 to 200 kbp, respectively, with 85 kbp of overlapping sequences (Figure 1C-D). An enhanced green fluorescent protein encoding reporter gene (Egfp) was inserted at the translation start site of Nppa in both BACs. Transgenic mouse lines were generated carrying the modified
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Figure 4. Postnatal expression of the Hprt-0.7rLacZ transgene compared to the -0.7rLacZ transgenic lines A and B. Whole mount β-galactosidase staining. Ventricular expression in the Hprt-0.7rLacZ transgene was not down-regulated after birth and remained active both in atria and ventricles (A, D). The expression in both -0.7rLacZ lines was inactivated in the atria as well as in the ventricles (B, E and C, F). Arrowhead in B indicates the small amount of ventricular activity present at day of birth in -0.7rLacZ line A. Arrowhead in F indicates the few atrial cells still expressing -0.7rLacZ in the adult atrium of line B. For abbreviations, see legend to Figure 2.

BAC clones. EGFP expression in five independent mouse lines carrying BAC337-Egfp showed similar spatio-temporal patterns, but different expression levels (Figure 5C-D, G, J). All lines showed ectopic expression in the neural tube (Figure 5C). In situ hybridization onsections showed robust expression of Egfp in the atria, but also ectopic expression in the sinus horns and in the mediastinal myocardium (Figure 5G), in a pattern similar to that of the -0.7rLacZ constructs. In the ventricles, the spatial pattern of Egfp expression resembled the endogenous Nppa pattern, restricted to the trabecules after E9.5 (Figure 5G, J). However, the observed ventricular fluorescence as well as the signal in in situ hybridization was very low when compared to the atria (Figure 5D, G). Three weeks after birth no ventricular fluorescence was detectable anymore (Figure 6D-F).

Two independent mouse lines carrying BAC336-Egfp showed a similar pattern and intensity of expression. Expression was restricted to the heart with abundant expression in the atria and ventricles (Figure 5A-B, F, I). The pre- and postnatal patterns of expression of Nppa and BAC336-Egfp were identical, including absence of expression from the mediastinal myocardium and sinus horns (Figure 5E-F, H-I) and downregulation of ventricular activity after birth (Figure 6A-C), indicating that this BAC clone contains all regulatory sequences involved in fetal spatio-temporal Nppa gene regulation.
Figure 5. Pre- and postnatal activity and expression patterns of BAC336-Egfp and BAC337-Egfp transgenes compared to endogenous Nppa. Mice carrying BAC336-Egfp showed heart-specific expression (A) with high activity both in the atria and the ventricles (B). Transgenic mice carrying BAC337-Egfp showed ectopic fluorescence outside the heart (C). Within the heart atrial fluorescence was abundant, but only minor activity was observed in the ventricles (D). In situ hybridization revealed that BAC336-Egfp mimics both pattern and activity of endogenous Nppa expression (E-F and H-I). In contrast, BAC337-Egfp in the ventricles recapitulated the pattern of endogenous Nppa (E, G and H, J), but was expressed at a very low level relative to atrial expression (G). BAC337-Egfp was ectopically active in the right and left sinus horns, the pulmonary vein, the mediastinal myocardium (arrow head) and the developing atrial septum (G). (K-L) qRT-PCR quantification of mRNA levels in E17.5 and adult hearts of BAC336-Egfp and BAC337-Egfp mice. (K) Ventricular activity of endogenous Nppa and BAC336-Egfp were approximately 5% of atrial activity before birth, and decreased to 1% of atrial activity after birth. In contrast, fetal ventricular BAC337-Egfp expression was less than 1% of atrial expression and after birth remained constant at this level. (L) Ratio of Egfp expression relative to endogenous Nppa expression in fetal atria and ventricles of two BAC336-Egfp and three BAC337-Egfp transgenic lines. In both BAC336-Egfp lines, the ratio of Egfp and Nppa expression in the atria and in the ventricles was similar, while in all BAC337-Egfp lines ventricular expression was only roughly 10% of atrial expression. Egfp and Nppa expression in the atria and the ventricles was normalized to Gapdh expression. Error bars indicate SEM. as, atrial septum. See legends to Figure 2 and 3 for other abbreviations.
To assess whether the BAC sequences provide correct fetal ventricular activity and perinatal down-regulation, the atrial and ventricular expression levels of Nppa and Egfp were quantified before and after birth (Figure 5K). At E17.5, ventricular expression of both Nppa and BAC336-Egfp were around 5% of atrial expression. After birth, both were down-regulated in the ventricles to 1% of atrial expression. In contrast, in E17.5 embryos carrying BAC337-Egfp, ventricular Egfp expression was less than 1% of atrial Egfp expression, and remained at this low level after birth. Subsequently, we calculated the ratios of Egfp and Nppa mRNA levels in E17.5 atria and ventricles of two BAC336- and three BAC337-Egfp transgenic lines (Figure 5L). In both BAC336-Egfp lines the ratio of the levels of Egfp and Nppa mRNA in the atria was similar to the ratio in the ventricles, whereas in all BAC337-Egfp lines the ratio of Egfp and Nppa mRNA in the ventricles was around 10% of the ratio found in the atria. These findings indicate that BAC337-Egfp lacks a ventricular enhancer that is mainly active before birth, locating this enhancer within the unique sequences of BAC336-Egfp (Figure 7).

**Distal regulatory sequences control the stress response of Nppa**

In the Hprt locus ventricular activity is maintained even in the adult heart. To test whether the 0.7 kbp proximal Nppa promoter in this transcriptionally favorable context is inducible in the failing ventricle in vivo, we subjected these mice to cardiomyopathy, using a transgenic model in which Gal4 is driven by the αMHC promoter (αMHC-Gal4) (Habets et al., 2003). All male Gal4-positive offspring develop a-specific dilated cardiomyopathy three weeks after birth, which is associated with strong induction of Nppa. Although endogenous Nppa was strongly induced in double transgenic mice, the Hprt-0.7rLacZ construct was not (Figure 6G). Subsequently we tested the 7 kbp fragment, which lacks the capacity for embryonic ventricular expression, but contains an NRSE implicated in induction of Nppa promoter fragments in heart failure (Kuwahara et al., 2003). This -3/+4mCre construct was not re-activated in the ventricles either (Figure 6G).

In contrast, mice carrying either BAC336-Egfp or BAC337-Egfp showed strong re-activation of Egfp in the failing ventricles (Figure 6A, D, G). In ventricles of BAC336-Egfp mice, Egfp was up-regulated 12-fold relative to its level in healthy littermates, which is similar to the up-regulation of endogenous Nppa. In mice carrying BAC337-Egfp, an exceptional 50-fold induction of Egfp over basic ventricular expression level was observed, the height of the induction being at least in part due to the low basic expression level of BAC337-Egfp in the healthy ventricle. Also transverse aortic constriction of mice carrying these BACs induced the expression of both Nppa and Egfp (Figure 6B, C, E, F; Supplementary Figure 3), demonstrating that the regulatory sequences in both BACs mediate the response of Nppa to pressure overload hypertrophy. These results show that both BAC clones contain the regulatory sequences required for the cardiac stress response of Nppa (Figure 7).
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Figure 6. BAC336-Egfp and BAC337-Egfp, but not the Hprt-0.7rlacZ and -3/+4Cre constructs, are reactivated in heart failure. (A, D) Heart failure induced by crossing the transgenic mice to the αMHC-Gal4 heart failure mouse (Habets et al., 2003). (B, E) Hypertrophy induced in adult mice using a pressure overload model (transverse aortic banding (TAC)). From the hearts carrying BAC337-Egfp (E) the atria were removed. (C, F) Midventricular transverse sections of hearts shown in B and E. (G) qRT-PCR quantification of ventricular mRNA levels of endogenous Nppa and the reporter genes LacZ, Cre and Egfp relative to Gapdh mRNA in control (black bars; mean ratio set to 1) and failing hearts (grey bars) in mice carrying the Hprt-0.7rlacZ, -3/+4mCre, BAC336-Egfp or BAC337-Egfp construct. In all four groups, the level of Nppa in failing hearts was significantly higher than in control hearts (P<0.05). In contrast, reporter gene expression in neither the Hprt-0.7rlacZ (n=4, P=0.3), nor the -3/+4mCre failing ventricles (n=3, P=0.9), showed induction. In mice carrying BAC336-Egfp (n=4, P<0.01) or BAC337-Egfp (n=3, P<0.05), we detected significant re-activation of Egfp in failing ventricles. Error bars indicate SEM. RG, reporter gene. See legend to Figure 2 for other abbreviations.
Discussion

Proximal Nppa promoter fragments lack regulatory functions for ventricular activity

In the developing heart, Nppa serves as a marker for differentiating atrial and ventricular working myocardium, whereas its expression is excluded from the sinus horns, atrioventricular canal, outflow tract and nodes of the conduction system (Zeller et al., 1987; Wharton et al., 1988; Christoffels et al., 2000). After birth, Nppa is expressed at very high levels in the atria (approximately 1% of total mRNA), whereas its expression in the ventricles is down-regulated (Bloch et al., 1986). Re-activation of Nppa expression is part of a highly conserved adaptive change in gene expression in response to hypertrophy and heart failure, serving both diagnostic and potentially therapeutic options. Because of these properties, Nppa has become a widely used model gene for studying gene regulation and monitoring phenotypic changes during cardiac disease (reviewed in (Levin et al., 1998; Schmitt et al., 2003; Houweling et al., 2005)). Proximal promoter fragments are frequently used as read-out tools for the activity of a wide range of transcriptional pathways that control patterning of the developing heart and gene regulation in healthy, hypertrophic or failing ventricular myocardium (reviewed in (Nemer and Nemer, 2001; Temsah and Nemer, 2005; Houweling et al., 2005)). It is therefore important to know which regulatory functions the proximal promoter contains. Promoter fragments ranging from -3.4 kbp to -0.5 kbp were reported to efficiently drive expression in the atria, and in several cases also in the ventricles of transgenic mice and Xenopus (Field, 1988; Seidman et al., 1991; Knowlton et al., 1995; Habets et al., 2002; Small and Krieg, 2003). Furthermore, analysis of promoter activity in transfected atrial and ventricular myocytes and non-cardiac cells has indicated that the Nppa promoter is organized in three modules, a basic cardiac promoter, a developmental- and atrial-specific module, and a ventricular enhancer located just upstream of the atrial module (Durocher et al., 1996; Durocher et al., 1998; Durocher and Nemer, 1998). These studies have led to the current view that the proximal Nppa promoter correctly drives atrial and fetal ventricular expression, and postnatal ventricular down-regulation.

Our current data reveal that the proximal promoter fragments lack important regulatory functions required for fetal ventricular activity. All promoter fragments analyzed were active in the atria and correctly inactive in the atroventricular canal and outflow tract of the developing heart, and in the nodes of the mature heart (Figure 7). The ‘atrial’ module in the context of the cTnl promoter was found to be largely sufficient to provide these characteristics (Figure 7; Supplementary Figure 1). These results provide in vivo support for previous studies showing that the activity of the promoter is mediated by T-box factors and Nkx2-5 that act on sites within this ‘atrial’ module (Bruneau et al., 2001; Hiroi et al., 2001; Habets et al., 2002). In contrast, the ventricular regulatory characteristics appeared to largely lack from proximal promoter fragments. Firstly, the fetal transmural pattern of expression of the Nppa gene was not recapitulated in transgenic mice. Secondly, while
ventricular activity was observed in the favorable genomic context of the \textit{Hprt} locus, in randomly integrated constructs it was weak, and absent in the context of larger \textit{Nppa} fragments (up to -11/+5 kbp). Thirdly, the activity of the 0.7 kbp promoter was not correctly down-regulated in the ventricles after birth. In the context of the \textit{Hprt} locus the \textit{Nppa} promoter fragment remained active, and in the randomly integrated constructs the

\textbf{Figure 7.} Overview of the activity of regulatory \textit{Nppa} sequences. Black lines depict the \textit{Nppa} locus, gray lines depict DNA fragments described in literature, and dark green lines fragments analyzed in this study. All fragments were studied in transgenic mice, except for the first three (\textsuperscript{a}) which were analyzed in cultured cardiomyocytes. act., ventricular activity; patt., ventricular pattern; HT, hypertrophy; Ref, reference; nd, not determined; na, not applicable; -, no expression; +, expression; +/-, expression in some lines; ++, high activity (comparable to endogenous \textit{Nppa}). See legend to Figure 2 for other abbreviations.
inactivation was non-specific, as also the atria lost expression. Finally, analysis of the BAC transgenic mice revealed that fetal ventricular expression requires two distinct distal sequences. The ventricular transmural pattern requires additional regulatory sequences located between -27 kbp to -11 kbp and/or +5 kbp to +58 kbp relative to the Nppa gene, whereas fetal ventricular activity is provided by a distinct strong fetal ventricular enhancer that we located more upstream between -141 kbp to -27 kbp (Figure 7). Therefore, the atrial module within the proximal promoter may provide only residual ventricular activity, which is unmasked in a favorable genomic context. Taken together, we conclude that the activity pattern of the Nppa proximal promoter region is very useful to study atrial gene activity and repression in the nodes of the conduction system, but does not represent a physiologically relevant read-out for ventricular gene regulation in the developing and adult heart.

**Distinct regulatory sequences and divergent pathways drive fetal activity and stress response**

The proximal promoter of Nppa is responsive to hypertrophic stimuli in some experimental settings (Shubeita et al., 1990; Ardati and Nemer, 1993; von Harsdorf et al., 1997), but does not respond to ventricular hypertrophy in transgenic mice in vivo (Knowlton et al., 1995). Our analysis confirms and extends these observations in a different heart disease model. Firstly, we tested a larger fragment that contains an NRSE which has been implicated in the hypertrophy response of Nppa. While this NRSE is sufficient to mediate a response to hypertrophic stimuli in vitro through NRSF (Kuwahara et al., 2003), our results indicate this is not the case in vivo. Secondly, in previous studies the postnatal expression and the inducibility of Nppa promoter fragments has been tested in mice carrying randomly integrated promoter constructs, which, according to our findings, may become down-regulated by postnatal silencing that is not specific to the ventricles. We circumvented this potential problem by testing the inducibility of the 0.7 kbp promoter in the context of the Hprt locus that stays transcriptionally accessible throughout life. Again, the transgene was found to be non-responsive in the cardiac disease model. Taken together, all data consistently indicate that the proximal promoter region lacks critical sequences that mediate the induction of Nppa, rendering the fragment completely non-responsive to cardiac disease in vivo.

Based on the assumption that the proximal promoter fragments drive fetal ventricular activity, previous studies indicated that re-activation of the fetal gene program during cardiac disease is regulated by pathways distinct from those that regulate prenatal ventricular activity (Knowlton et al., 1995). As discussed above, our analysis shows that also the fetal ventricular activity functions are lacking from the proximal promoter fragments. Therefore, the previous conclusion regarding the divergent pathways was premature.

The analysis of two BAC transgenic mice revealed that ventricular Nppa regulation requires a multi-partite ventricular regulatory module. This module consists of the proximal 0.25 kbp ‘atrial’ module driving residual ventricular expression, and at least two additional distal sequences (Figure 7). Whereas BAC337-Egfp provided the correct fetal transmural
ventricular pattern, and was appropriately re-activated during disease, its ventricular activity before birth was similarly low as in the adult heart, indicating that BAC337 lacks a ventricular enhancer inducing activity before birth. In contrast, BAC336-Egfp activity mimicked all aspects of ventricular Nppa regulation, including fetal transmural pattern, abundant fetal activity, perinatal down-regulation, and re-activation in hypertrophy and in heart failure. These results indicate that shared sequences, residing within both BAC clones, may regulate the typical fetal transmural (trabecular) ventricular pattern and re-activation in cardiac disease (Figure 7). Importantly, this analysis also revealed that the sequences required for activity in the fetal ventricle are located in a different genomic region, more than -27 kbp upstream (Figure 7). Therefore, the re-activation in the failing ventricle and the fetal ventricular activity are regulated by distinct distally located sequences, and, consequently, divergent pathways converging on these sequences. Further mapping will be required to locate the discrete cis-elements and factors that mediate these respective activities.

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References


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Supplementary data

Supplementary materials and methods

Transgenic mice

Hprt-0.7rLacZ
Of the targeted Hprt-0.7rLacZ two independent transgenic lines were established in which we verified a single integration event at the Hprt locus by Southern blot. Both lines showed identical expression patterns.

-3mCre
To generate the -3mCre construct we truncated the -3/+4mCre construct upstream of the NRSE, by cloning the human growth hormone polyadenylation signal into the PvuI site located in the third exon of Nppa.

Bac336-Egfp and BAC337-Egfp
The two-step BAC modification protocol previously described by Shiaoching Gong and Nathaniel Heinz (Gong et al., 2002), consists of two homologous recombination steps. After both the co-integration and the resolution step correct recombination was verified by Southern blot using a hybridization probe against Egfp. The BAC-Egfp constructs were purified using a CsCl gradient following a protocol also kindly provided by Shiaoching Gong and Nathaniel Heinz.

-11/+5mEgfp
To generate the -11/+5mEgfp construct, we started with the modified BAC336-Egfp construct. Using conventional restriction enzymes we cloned a fragment reaching from the first natural Sal1 site at -11 kbp upstream of Nppa, to the first natural Xho1 site at +5 kbp downstream of Nppa.

Heart failure mouse model
Mice carrying the Hprt-0.7rLacZ or -3/+4mCre construct, or one of the modified BAC-Egfp clones, were crossed with the αMHC-Gal4 heart failure mouse previously described (Habets er al., 2003). Gal4 positive and double positive male offspring developed heart failure three
weeks after birth. When dyspnoea was diagnosed, mice were terminated. On postmortal examination all Gal4 positive male mice showed an enlarged heart and a swollen, yellowish liver, signs of heart failure with congestion. Atria and ventricles of double positive hearts and of Cre, LacZ or Egfp single positive hearts of littermates were separated and RNA was isolated using the Nucleospin RNA II kit (Macherey-Nagel, Düren, Germany) following the protocol of the manufacturer. First strand cDNA was synthesized using an optimized reverse transcription protocol (Lekanne Deprez et al., 2002).

**Aortic banding**

Transverse aortic constriction (TAC) was performed in 3- to 4-month-old mice, carrying the -3/+4mCre construct, or one of the modified BAC-Egfp clones. The aorta was subjected to a defined 25-gauge constriction between the first and second truncus of the aortic arch as described (Rockman et al., 1991). Aged-matched unbanded animals were used as controls. When discomfort (dyspnoe, decreasing mobility) was diagnosed between the 2nd and 3rd week after TAC, mice were terminated and heart tissue was collected. On postmortal examination all mice included in the qRT-PCR experiment had an enlarged heart and/or a stiffened, more solid appearance. RNA isolation and cDNA synthesis was performed as described above.

Animal care was in accordance with national and international guidelines.

**Quantitative real time PCR**

Quantitative real time PCR was performed using a LightCycler Real-Time PCR system (Roche Diagnostics, Almere, The Netherlands). The relative start concentration \( N(0) \) was calculated using the following equation: \( N(0) = 10^{[\log(\text{threshold})–Ct(\text{mean Eff})]} \). Values were normalized to Gapdh expression levels.

**Non-radioactive in situ hybridisation and β-galactosidase activity detection**

Whole mount in situ hybridization, in situ hybridization on sections and whole mount and cryosection β-galactosidase activity staining were performed as described previously (Moorman et al., 2001; Franco et al., 2001).
Supplementary figures

**Supplementary Fig. 1.** A 0.25 kbp ‘atrial’ regulatory module correctly represses *Nppa* in the AVC and drives atrial and residual ventricular expression. (A) A 356 bp *cTnI* promoter fragment is always expressed in the atrioventricular canal and only shows limited expression in the atria and ventricles. (B) As previously shown (Habets et al., 2002), a fragment of -638/-138 bp of the rat *Nppa* promoter is able to impose inhibition of expression upon the *cTnI* fragment in the atrioventricular canal, while increase of expression is observed in the atria and to a lesser extent in the ventricles. (C) All elements required to inhibit expression in the atrioventricular canal and to drive the atrial and ventricular expression reside within a 0.25 kbp module of the *Nppa* promoter. ra, right atrium; la, left atrium; rv, right ventricle; lv, left ventricle; avc, atrioventricular canal; oft, outflow tract.

**Supplementary Fig. 2.** Correct ventricular expression requires distal regulatory sequences that reside outside a -11 to +5 kbp promoter fragment. (A) As previously shown (de Lange et al., 2003), extension of the promoter fragment to 7 kbp (-3/+4mCre) diminishes ventricular activity. This lack of activity is not caused by a specific ventricular repressor located either up- or downstream of the 0.7 kb proximal promoter fragment as indicated by the lack of ventricular activity in the -3mCre (B) and the +4mLacZ transgenes (C). (D) A mouse -0.7mLacZ promoter fragment behaves similarly as the rat -0.7rLacZ promoter fragment, with activity in both the atria and the ventricles and ectopic expression in the sinus horns. (E) A promoter fragment of 16 kbp (-11/+5mEgfp) does not drive any activity in the ventricles either. Arrowheads point at the sinus horns. See legend to supplementary Fig. 1 for other abbreviations.
Supplementary Fig. 3. Induction of Nppa, BAC336-Egfp and BAC337-Egfp, but not the -3/+4Cre construct, in mice after induction of hypertrophy by transverse aortic constriction (TAC). In all mice carrying the BAC336-Egfp (n=3) or the BAC337-Egfp (n=3) construct, green fluorescent protein was visibly upregulated upon pressure overload. This finding was confirmed by quantitative RT-PCR. All groups >/=3. Error bars indicate SEM. RG, reporter gene.

References supplementary data


Interplay between Tbx20, Tbx2/Tbx3 and Bmp/Smad-signaling controls compartmentalization of the developing heart tube into working chambers and the atrioventricular canal

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Abstract

The generation of efficient unidirectional blood flows in the mammalian heart relies on the functional compartmentalization into chambers separated by valves and septa. Here, we report on a genetic network involving the T-box transcription factor genes Tbx20, Tbx2 and Tbx3 and Bmp/Smad-signaling that coordinates the formation of the working chambers and the atrioventricular (AV) canal, from which the AV conduction system and the mesenchymal cushions, primordia of valves and septa, arise during cardiac development. We show that Tbx2 and its close relative Tbx3 are redundantly required and individually sufficient to specify AV myocardium, induce formation of the AV cushions and suppress chamber differentiation. Tbx20 is required for chamber formation independently from Tbx2, but suppresses Tbx2 in the developing chambers, thereby localizing its activity to the AV canal. We identified a Bmp/Smad-dependent enhancer conferring AV canal restricted expression and Tbx20-dependent chamber suppression of Tbx2 in vivo. Unexpectedly, Tbx20 does not repress expression of Tbx2 and other genes by binding to DNA-elements, but attenuates Bmp/Smad-dependent activation by binding Smad1 and 5 and sequestering them from Smad4. Our findings suggest that opposing regulation of Bmp-signaling by Tbx20 and Tbx2 may underlie specification of the chambers and the AV canal, respectively.
Introduction

The complex multi-chambered heart of vertebrates arises from a simple tubular structure through a coordinated program of cellular differentiation and proliferation, and tissue morphogenesis. Elongation of this simple tube is supported by recruitment of precursor cells and differentiation into cardiomyocytes at the two poles. Highly localized processes of further myocardial differentiation and increased proliferation within the growing heart tube mediate the local out-bulging of the atrial and ventricular chambers. Regions separating and bordering the developing chambers retain low proliferation rates and slow impulse conduction and resist differentiation in a working type of myocardium, resulting in the generation of primitive morphological constrictions, the AV canal (AVC) and outflow tract, and a delay in AV conduction (Moorman and Christoffels 2003). The primary AV myocardium, from which the definitive AV node derives, induces the overlying endocardium to undergo an epithelial-mesenchymal transition (EMT) and to invade the cardiac jelly, an extracellular matrix that is deposited by the primary myocardium. These mesenchymal cushions are subsequently remodeled into thin valve leaflets and components of the septa (Person et al. 2005) that ensure structural and functional compartmentalization of the heart.

Formation of the initial heart tube from the cardiac crescent and further elongation of the tube from mesodermal progenitor pools, requires Bmp-signaling, which induces the differentiation into cardiomyocytes (Klaus et al. 2007; Yang et al. 2006). Bmp-signaling regulates expression of an evolutionary conserved network of transcription factor genes that drives the development of chambers and the differentiation of working myocardium therein (Klaus et al. 2007; Prall et al. 2007). Ablation of any member of the network, including Nkx2-5, Gata and MeF2 factors, and the T-box transcription factors Tbx5 and Tbx20 causes major heart defects and early developmental arrest. Mice homozygous mutant for Nkx2-5 and Tbx20 establish a heart tube with a primary myocardial phenotype but fail to undergo looping morphogenesis and to initiate chamber formation (Cai et al. 2005; Singh et al. 2005; Stennard et al. 2005; Takeuchi et al. 2005; Prall et al. 2007). Tbx5 acts independently of Tbx20 and maintains posterior domains of the heart (Bruneau et al. 2001). Both Tbx20 and Tbx5 synergize on the biochemical level with other members of the conserved network, Nkx2-5 and Gata4, to activate expression of chamber specific genes such as Nppa and Cx40 (for a review see Hoogaars et al. 2007a).

Formation of the AVC again relies on Bmp-signaling (Sugi et al. 2004; Ma et al. 2005) and is elaborated and stabilized by other pathways (Rutenberg et al. 2006; Kokubo et al. 2007). Transcriptional repressor Tbx2 acts downstream of Bmp-signaling (Yamada et al. 2000) and regionally inhibits a chamber myocardial gene program in the AV canal by competing with activating T-box proteins such as Tbx5 for binding to conserved T-box binding elements (TBE)s in promoters of chamber specific genes (Habets et al. 2002; Christoffels et al. 2004b; Harrelson et al. 2004; Hoogaars et al. 2004; Chi et al. 2008). Tbx3 is genetically and functionally related to Tbx2, and suppresses chamber differentiation of the
sinus node, the AV bundle and bundle branches (Hoogaars et al. 2007b; Mommersteeg et al. 2007; Bakker et al. 2008). Tbx2 and Tbx3 expression overlaps in the AV canal, suggesting that functional redundancy has prevented a full appreciation of their role in the development of this tissue to date (Ribeiro et al. 2007; Bakker et al. 2008; Mesbah et al. 2008).

Although important signaling and transcriptional modules involved in the establishment of the cardiac components have been identified (Olson 2006; Dunwoodie 2007), the genetic hierarchies and the spatial and temporal interplay of these pathways have remained elusive. How for example Bmp-signaling, after inducing cardiogenesis, is redeployed to locally activate Tbx2 expression and AV canal formation has remained unclear, as has the role of core cardiac transcription factors in orchestrating these morphogenetic processes. The observation that Tbx2 is ectopically up-regulated in the entire embryonic heart tube of Tbx20-deficient embryos, possibly causing the block in chamber differentiation in Tbx20-deficient hearts, has suggested that Tbx20 is required to repress Tbx2 for the progression to a multi-chambered heart (Cai et al. 2005; Singh et al. 2005; Stennard et al. 2005). Here, we present genetic experiments in the mouse that further decipher the molecular pathways that underlie localized formation of the chambers and the AV canal. We show that Tbx20 plays a dual role in compartmentalization of the heart. It stimulates chamber differentiation independently from Tbx2 and simultaneously suppresses Bmp/Smad-dependent activation of Tbx2 in the heart tube by a previously unrecognized DNA binding-independent mechanism, thereby restricting its expression to the prospective AVC region. Tbx2, together with Tbx3, subsequently impose the AV canal phenotype on this region of the heart tube. Together, our data provide insight into the hierarchical mechanisms underlying the spatial delimitation of a broadly active signaling pathway and its redeployment to compartmentalize the heart.

**Materials and methods**

**Mice and genotyping**

Mice carrying a null allele of Tbx20 (Tbx20^tm1Akis, synonyms: Tbx20^−, Tbx20^lacZ) (Singh et al. 2005), Tbx2 (Tbx2^tm1.1(cre)Vmc, synonyms: Tbx2^−, Tbx2^Cre) and/or Tbx3 (Tbx3^tm1.1(cre)Vmc, synonyms: Tbx3^−, Tbx3^Cre) (Hoogaars et al. 2007b) were maintained on an outbred (NMRI) background. The Tbx2^Cre transgenic line, which harbors a Cre gene at the translation start site and from which the Pgk-neo cassette was removed, is a null allele that will be described elsewhere. For the generation of mutant embryos, heterozygous mice were intercrossed. For the generation of double mutant embryos, double heterozygous mice were intercrossed. For timed pregnancies, vaginal plugs were checked in the morning after mating, noon was taken as embryonic day (E) 0.5. Embryos were harvested in PBS, fixed in 4% paraformaldehyde overnight and stored in 100% methanol at −20°C before further use. Wild-type littermates were used as controls. CAG-CAT-TBX3 (CT) and Nppa-Cre4 (Cre4) transgenic mice have
been described (Hoogaars et al. 2007b). Double-transgenic mice conditionally expressing TBX3 in the atria were generated by crossing CT mice with Cre4 mice. Genomic DNA prepared from yolk sacs or tail biopsies was used for genotyping by PCR (Details are available on request).

An expanded Materials and Methods section is available in the online data supplement and included at the end of this chapter.

Results

Tbx20 is required for chamber formation, whereas Tbx2 stimulates cardiac jelly formation

Previous analysis has shown that ectopic expression of Tbx2 in the simple heart tube leads to arrest of cardiogenesis (Christoffels et al. 2004b). To test the hypothesis that ectopic Tbx2 expression is responsible for the observed block in chamber differentiation in Tbx20-deficient hearts (Cai et al. 2005; Singh et al. 2005; Stennard et al. 2005), we generated embryos with combined deficiencies of Tbx2 and Tbx20 by interbreeding double heterozygous animals. Similar to Tbx20 single mutants, Tbx20lacZ/lacZ;Tbx2Cre/Cre embryos were severely growth retarded and died at E10.5 due to hemodynamic failure (data not shown). At E9.5, when cardiac chambers were clearly delineated in wild-type embryos, Tbx20 single mutant and Tbx20/Tbx2 double mutant embryos featured a simple heart tube (Fig. 1A-C). On histological stainings of sagittal sections both myocardium and endocardium appeared homogenously thin throughout the Tbx20lacZ/lacZ;Tbx2Cre/Cre heart tube that sometimes appeared dilated. This is in contrast to Tbx20lacZ/lacZ hearts, where a thick layer of cardiac jelly filled the space between myocardium and endocardium. Cardiac jelly production was not associated with endocardial EMT and cushion formation in the Tbx20lacZ/lacZ linear heart tube as revealed by histological inspection as well as expression analysis of cushion markers Sox9 (Akiyama et al. 2004) and Msx2 (Abdelwahid et al. 2001) (Fig. 1D-F and data not shown). Molecular analysis using markers with differential expression along the linear heart tube (Tbx5, Myh7) (Bruneau et al. 1999;Christoffels et al. 2000) confirmed that anterior-posterior patterning occurred normally in Tbx20lacZ/lacZ;Tbx2Cre/Cre hearts (Fig. 1G-L). Chamber myocardium, however, was not formed, as shown by absence of expression of Nppa (Anf) and Cited1 (Fig. 1M-R). Bmp2 that is expressed in the primary myocardium of the AV canal in the wild-type (Ma et al. 2005), was found throughout the tubular heart of single and double mutant embryos (Fig. 1S-U). In summary, loss of Tbx2 does not rescue the Tbx20-deficient heart from developmental arrest at the linear heart tube stage, but results in loss of cardiac jelly formation. These data show that Tbx20 is a critical factor in cardiac chamber development, and that Tbx2 is sufficient to induce the first step in cushion formation, the production of a thick layer of cardiac jelly in the developing heart.
Figure 1. Loss of Tbx2 does not rescue cardiac arrest of Tbx20-deficient mice. Comparative analysis of wild-type, Tbx20+/− and Tbx20+/−Tbx2+/− embryos for cardiac morphology, histology and molecular marker expression at E9.5. (A-C) Left lateral views of whole E9.5 embryos reveal growth retardation and linear heart tube phenotype in Tbx20/Tbx2 double mutants similar to Tbx20-deficient embryos. Boxes mark the heart regions to be magnified in the following images. (D-F) Histological analysis of sagittal sections by hematoxylin and eosin (H&E) staining uncovers loss of cardiac jelly and cushion tissue (asterisk) in the Tbx20/Tbx2-deficient linear heart tube that appears as a slightly inflated sac. (G-U) In situ hybridization analysis of marker gene expression in whole embryos with probes as indicated. avc, atrioventricular canal; ec, endocardium; la, left atrium; lv, left ventricle; mc, myocardium; oft, outflow tract; pv, primitive ventricle; rv, right ventricle.
Tbx2 and Tbx3 are cooperatively required for myocardial patterning and formation of cushion mesenchyme in the AV canal

Previous analyses indicated that Tbx2 is necessary and sufficient to suppress chamber gene expression in the AV canal (Christoffels et al. 2004b; Harrelson et al. 2004). However, in our Tbx2 loss-of-function mouse AV canal formation at E9.5 was grossly normal, and Nppa and other chamber markers were not ectopically expressed in the AV canal (Fig. 2F,J,N).

![Figure 2](image-url)
However, co-expression with the close family member *Tbx3* in the AVC myocardium argues that functional redundancy precludes full appreciation of *Tbx2/Tbx3* requirement in this region. We therefore aimed to generate mice double mutant for *Tbx2* and *Tbx3*. Mice double heterozygous for *Tbx2* and *Tbx3* null alleles are afflicted with craniofacial defects that lead to early postnatal lethality (unpubl. observ.). Breeding onto an NMRI wild-type background partially rescued this phenotype so that double heterozygous mice could be interbred. Since pregnancies were badly maintained in these mothers, we obtained only a total of four viable *Tbx2/Tbx3* double homozygous embryos from few litters at E9.5. These embryos appeared slightly retarded in their development, most likely due to arising hemodynamic failure. Morphologically, the constriction between the left ventricle and the atrium in the wild-type was largely absent (Fig. 2C). Histologically, the AV canal appeared as an inflated tube with little investment of cardiac jelly, and, in contrast to the other genotypes, complete absence of cushion tissue (Fig. 2G). Markers of chamber myocardium (*Nppa, Gja5 (Cx40)*) (Christoffels et al. 2004a) were expanded into this region, whereas markers for primary myocardium of the AV canal, *Bmp2* (Ma et al. 2005) and *Cre* from the mutant alleles reflecting endogenous *Tbx2* and *Tbx3* expression, were present but reduced in their expression levels and spatial extent (Fig. 2I-X). A similar phenotype was observed in compound mutants with loss of three functional alleles of *Tbx2* and *Tbx3* (Suppl. Fig. 1), whereas *Tbx2* and *Tbx3* single mutants showed normal cushion formation and proper chamber gene repression in the AV canal (Fig. 2). Hence, *Tbx2* and *Tbx3* are required in a redundant fashion to maintain, expand and differentiate the primary myocardium of the AV canal, and to induce the formation of cushion tissue from the endocardium in this region.

**Tbx3 regulates the AV myocardial gene program**

The difficulty in obtaining a sufficiently high number of *Tbx2*\(^{Cre/Cre} \cdot Tbx3^{Cre/Cre}\) embryos for a more detailed analysis of the molecular consequences of combined loss of *Tbx2* and *Tbx3*, prompted us to use a complimentary gain-of-function approach to further elucidate the role of these factors in AV canal development. We used ectopic expression of *Tbx3* in the atrial myocardium that we have previously shown to be powerful in achieving molecular insight into cardiac *Tbx3* function (Hoogaars et al. 2007b). Nppa-Cre4 (*Cre4*) mice were crossed with *CT* mice to obtain permanent activation of *Tbx3* in atrial chamber myocardium from E10.5 onwards in double transgenic *Cre4;CT* mice (Hoogaars et al. 2007b). Micro-array analysis comparing the transcriptome of atria of adult *Cre4;CT* and *Cre4* males confirmed earlier findings of strong reduction in expression of the known chamber genes *Nppa, Smpx (Chisel), Scn5a (Nav1.5), Kcnj3 (Kir3.1), Gja1 (Cx43) and Gja5 (Cx40)* in *Tbx3*-expressing atria of *Cre4;CT* mice (Hoogaars et al. 2007b), and additionally revealed down-regulation of *Ckm, Nppb, Bmp10, Ednra* and *Aldh1b1* (Suppl. Table 1). qRT-PCR and *in situ* hybridization analysis confirmed normal chamber restricted expression and *Tbx3*-mediated atrial repression of *Ckm* (Wessels et al. 1990), *Nppb* (Houweling et al. 2005), *Aldh1b1* and *Ednra* (Clouthier et
Figure 3. Myocardial Tbx3 expression induces endocardial mesenchyme formation and nodal gene expression. Sections of E17.5 atria of CT and Cre4;CT mice were probed for expression of indicated genes. Note the reduced complexity and smoothness of the pectinated muscle structure in atria of Cre4;CT fetuses. Black arrow head indicates sinus node (san), white arrow head the myocardium in which Cacna2d2, transgenic TBX3 and Bmp2 expression was seen to be induced in Cre4;CT atria. Red arrow heads depict the thick endocardial mesenchymal layer that forms in Cre4;CT atria, that is devoid of myocardial gene expression (cTnI), but expresses Fbln2 and Tgfβ2. Black bar, 100µm.

Previously, we observed that nodal conduction system genes Hcn1, Hcn4, Cx30.2, Cav3.1 and Lbh are induced in Cre4;CT mice (Hoogaars et al. 2007b), indicating that Tbx3 stimulates the nodal gene program. Micro-array analysis and subsequent validation by qRT-
PCR and/or in situ hybridization revealed induction of additional genes in Cre4;CT atria including Cx45, Itpr1, Slco3A1, Id2, Cacna2d2, and Hnt (Fig. 4). These genes are enriched in the conduction system components, including the AV node, and associated with (Slco3A1, Cacna2d2, Hnt) or involved in the formation (Id2) or function (Cx45, Itpr1) of the conduction system (Gorza et al. 1993; Coppen et al. 1999; Kreuzberg et al. 2005; Mery et al. 2005; Moskowitz et al. 2007) (Fig. 3, 4A,B). Together, these data indicate that Tbx3 is able to suppress a large number of chamber genes that are reduced in the AV canal and conduction system, and to induce AV canal-enriched genes implicated in conduction system specification or function.

**Tbx3 induces sub-endocardial mesenchyme formation and Bmp-signaling**

Histological analysis of Cre4;CT atria and controls at E17.5 revealed that the pectinated muscles, trabecule-like structures in the atrial appendages, were abnormally thick, smooth and uncomplicated and that a thick sub-endocardial layer of cells had formed (Fig. 3). This tissue expressed mesenchymal marker genes including Acta2/αSMA, Fbln2, Cspg2/Versican and Lumican (Fig. 3, 4A, Suppl. Table 1) that are associated with AV cushions. In addition, micro-array analysis and in situ hybridization revealed strong activation of components of the Tgfβ-, Bmp-, Fgf- and Wnt-signaling pathways that have been functionally implicated with AV cushion and valve formation (Armstrong and Bischoff 2004; Person et al. 2005). Notably, we detected induction of Bmp2 expression in the myocardium, and of Bmp4 in endocardial layers of Cre4;CT mice (Fig. 3, 4A). Since Bmp2 expression in the AV canal myocardium is both required and sufficient to induce cushion formation (Yamagishi et al. 1999; Sugi et al. 2004; Ma et al. 2005), Bmp2 is likely to be a pivotal downstream mediator of cushion induction by Tbx3. Tgfβ2, a Bmp2 target in this tissue that is required for cushion formation (Camenisch et al. 2002; Mercado-Pimentel and Runyan 2007) was upregulated in the endocardium (Fig. 3, 4A). In addition, qRT-PCR and in situ hybridization analyses confirmed expression of genes in the endocardial mesenchymal layer of Cre4;CT mice (Twist1, Mx1, Meox1, Sox9, Id3 and Smad6) (Fig. 4, Suppl. Fig. 2 and 3, Suppl. Table 1), whose expression and functional relevance in EMT and cushion formation in the AV canal have been reported (Galvin et al. 2000; Akiyama et al. 2004; Ma et al. 2005; Lincoln et al. 2006; Chen et al. 2008; Shelton and Yutzey 2008). Furthermore, expression of Fgfr2, and of Wnt antagonists Fz8b, sFRP2 and Nkd2 were up-regulated in atria of Cre4;CT mice, compatible with the known role for Fgf- and Wnt-signaling in cushion and valve formation (Gitler et al. 2003; Hurlstone et al. 2003; Sugi et al. 2003; Person et al. 2005) (Suppl. Fig. 2, 3). Pkd2 is normally expressed in the developing valves and was induced in atrial mesenchyme of Cre4;CT mice. (Fig. 4A). In human and mouse, mutations of Pkd2 result in valve abnormalities (Wu et al. 2000; Stypmann et al. 2007). Finally, we identified additional 47 induced genes in the microarray data whose specific expression in the fetal AV valvular mesenchyme was reported by Genepaint (http://www.genepaint.org/) (Suppl. Table 2).
In summary, these data show that ectopic myocardial expression of \textit{Tbx3} is sufficient to induce the mesenchymal transition of endocardial cells and the formation of AV cushion tissues. Hence, Tbx3 collectively regulates the genetic circuits of signaling pathways and transcriptional activities mediating these processes.

\textbf{Figure 4.} Tbx3 suppresses chamber-specific genes and induces genes involved in cardiac cushion formation and conduction system function. (A) qRT-PCR analysis of left atria of Cre4-CT double transgenic mice compared to CT control mice. Expression levels in CT atria were set to 1. Expression of chamber-specific genes was reduced, whereas genes associated with conduction system were induced (nodal/channel). Genes encoding signaling proteins and transcription factors involved in cardiac cushion formation and valve development were induced in atria of Cre4-CT mice. Error bars represent SD (n=4 per group). *P<0.05. (B) Cross-sections of E12.5 mouse hearts showing expression of genes up-regulated in Cre4;CT atria (micro-array) in the developing AV canal and AV cushions and valves.
**Regulation of cardiac Tbx2 expression by Bmp/Smad-signaling and Tbx20**

To gain insight into the molecular pathways that confine Tbx2 expression to the developing AVC, we functionally examined the Tbx2 regulatory region. We first tested a 6 kbp genomic fragment previously shown to mediate AVC expression (Kokubo et al. 2007), and found that Eyfp reporter gene expression driven by this fragment in transgenic animals recapitulated endogenous cardiac expression of Tbx2 in the AV canal and outflow tract in E9.5 to E11.5 wild-type embryos. Since this fragment also conferred ectopic expression in Tbx20-deficient hearts (Fig. 5A-G), all elements required for control of cardiac Tbx2 expression reside within this genomic DNA fragment that contains several phylogenetically conserved regions (Fig. 5A, Suppl. Fig. 4). Deletion analysis of this fragment showed that the previously identified TBE recognized by Tbx20 (Cai et al. 2005) is neither required for AV canal and outflow tract-specific expression nor Tbx20-mediated repression. We identified a 0.9 kbp genomic fragment located 2.3 kbp upstream of the transcriptional start site, that in combination with a minimal promoter piece of 0.6 kbp (-1.5mTbx2) recapitulated cardiac expression of Tbx2 in wild-type and up-regulation in Tbx20-deficient embryos faithfully (Fig. 5H-J, Suppl. Fig. 5). Recent reports have pinpointed the relevance of conserved Foxn and additional TBE sites in the zebrafish tbx2b promoter for activation of the gene in the AV canal (Chi et al. 2008). Moreover, b-catenin has been shown to regulate Tbx3, the T-box factor most closely related to Tbx2, in cancer cells by direct binding to a Lef/Tcf site (Renard et al. 2007). A 1.5 kbp genomic fragment deprived of these conserved sites still drove reporter gene expression to the AV canal and outflow tract, negating a role for T-box factors, Lef/Tcf proteins and Fox transcription factors in regulating cardiac Tbx2 expression on the DNA level in the mouse (Fig. 5A, Suppl. Fig. 5). However, this -1.3mTbx2 genomic fragment contained a large number of putative Smad binding sites (SBE), supporting a role of Bmp-signaling in the regulation of cardiac Tbx2 expression (Suppl. Fig. 4). Taken together, this analysis indicates that a small SBE-containing enhancer in Tbx2 is sufficient to drive AV canal expression in vivo.

We switched to in vitro reporter assays in NIH3T3 cells to further investigate the precise role of Tbx20 in conjunction with Bmp/Smad-signaling in cardiac Tbx2 expression (Fig. 5K,L). We used expression constructs for full-length mouse proteins for Smad5, the constitutively active Bmp receptor Alk3 (Alk3CA) (Wessely et al. 2001) and HA-tagged Tbx20, and a reporter plasmid containing the luciferase reporter gene downstream of the 6 kbp Tbx2 genomic fragment in these transactivation assays. We observed a strong activation of reporter gene activity upon co-expression of Smad5 and Alk3CA and a reduction to basal activity in the presence of increasing concentrations of Tbx20 (Fig. 5L). Deletion analysis of the Tbx2 promoter fragment revealed the requirement of a 0.9 kbp Nhelli/AflIII fragment for Bmp/Smad-dependent activation of the promoter (Fig. 5K, Suppl. Fig. 6). Since this fragment was found to be sufficient to confer AV canal/outflow tract expression of Eyfp in vivo, Bmp/Smad-signaling is likely to activate cardiac Tbx2 expression in vivo. Removal of previously identified TBEs (Cai et al. 2005) did not affect the repression activity of Tbx20 on...
the promoter (Fig. 5K, Suppl. Fig. 6). This may suggest the presence of cryptic DNA binding sites for Tbx20. Alternatively, repression by Tbx20 may not be mediated by DNA binding but by protein interaction. To test the latter, we constructed point mutants of Tbx20 that do not exert specific DNA binding any more (Suppl. Fig. 7B,C). Unexpectedly, these mutant Tbx20 proteins still repressed transactivation of the 6 kbp Tbx2 fragment by Bmp/Smad- signaling.

**Figure 5.** Bmp/Smad-signaling controls cardiac expression of Tbx2. In vivo and in vitro reporter analyses of the control of cardiac Tbx2 expression. (A) Phylogenetic analysis of a 6 kbp genomic region upstream and around the Tbx2 transcription start site for conserved sequences. The location of conserved T-box binding elements (TBE) and Foxn sites is marked by boxes. Deletion constructs of the 6 kbp Tbx2 genomic fragment used in vivo to drive Eyfp reporter expression. Presence (+) and absence (-) of Eyfp expression from these deletion constructs in the atrium (A), atrioventricular canal (AVC), ventricle (V) and outflow tract (OFT) in transgenic mouse embryos. Numbers indicate cardiac expression of Eyfp and the number of transgenic embryos analyzed (Card. expr/tg) and extracardiac expression domains detected in the same embryos (Extra card. exp/tg). A 0.9 kbp NheI/AflII fragment from the Tbx2 locus contains elements sufficient and required to drive reporter gene activity in the regions of primary myocardium. (B-J) Comparative analysis of Tbx2 mRNA expression (B-D), Eyfp activity from the 6.0 kbp Tbx2 genomic region in transgenic embryos (E-G), and Eyfp activity from the 1.5 kbp Tbx2 genomic region in transgenic embryos (H-J) in E9.5 wild-type embryos (B,E,H, arrows point to the AVC), E10.5 hearts (C,F,I), and E9.5 Tbx20-deficient embryos (D,G), arrows point to mutant linear heart tube. (K) In vitro reporter assays to detect transcriptional activation of a luciferase reporter from Tbx2 genomic fragments. Plasmids encoding constitutively active Bmp receptor Alk3, full length Smad5 protein, full length Tbx20 protein and the luciferase reporter construct were co-transfected in NIH3T3 cells and luciferase activity determined and normalized as fold over the reporter alone. Presence (+) and absence of activation (-) by Bmp/Smad-signaling and repression by Tbx20 is listed for the constructs tested. (L), Luciferase reporter assay for the 6 kbp Tbx2 genomic fragment. Numbers indicate µg of plasmids for the reporter -6.0mTbx2-luc, and the expression plasmids for Alk3CA, Smad5, Tbx20 and the non-DNA binding Tbx20 protein (Tbx20 mut) co-transfected into NIH3T3 cells. Reporter activation upon co-transfection of expression constructs for Alk3CA and Smad5 is inhibited by co-expression of Tbx20, independent of its ability to bind to DNA.
both in presence and absence of transfected Smad5 in co-transfection experiments (Fig. 5L). Repression achieved by the DNA-binding deficient mutant of Tbx20 was lower than with the wild-type protein suggesting the co-existence of DNA-binding dependent and independent mechanisms for Tbx20 protein in the regulation of Tbx2 (Fig. 5L). Together, our analyses of the Tbx2 promoter argue that temporal and spatial confinement of cardiac Tbx2 expression is achieved by Tbx20-dependent repression of Bmp/Smad-mediated transcriptional activation.

**Figure 6.** Tbx20 inhibits Bmp/Smad-mediated transcriptional activation. (A-D) Luciferase reporter assays on a 0.7 kbp Msx2 genomic fragment (A), a 91 bp array of Smad bindings sites derived from the Id1 promoter (B,C), and the 0.7 kbp Nppa promoter fragment (D). Numbers indicate µg of plasmids for the luciferase reporter plasmids, and the expression plasmids for Alk3CA, Smad5, Tbx20 and the non-DNA binding Tbx20 proteins Tbx20mut1 (L126R), Tbx20mut2 (L127R) and Tbx20mut (LL126,127RR) co-transfected into NIH3T3 cells. Wild-type as well as the DNA-binding dead mutant Tbx20 protein inhibit dose-dependently activation by Bmp/Smad whereas the Tbx20 protein activates the Bmp/Smad-independent Nppa promoter.

**Tbx20 inhibits transcriptional activation of Bmp/Smad-dependent promoters in a DNA-independent manner**

We wondered whether the repressive effect of Tbx20 on Bmp/Smad-dependent transcriptional activation might be of a more general nature. We tested minimal fragments of Msx2 and Id1 promoters known to be activated by Smad binding in transactivation experiments in NIH3T3 cells (Brugger et al. 2004; Monteiro et al. 2004) (Fig. 6A-C). Both Tbx20 wild-type and Tbx20 DNA-binding mutant proteins repressed Bmp/Smad-mediated activation of Msx2 and Id1 promoter in a dose-dependent manner. Repression was independent from the addition of
exogenous Smad5 since it was also observed when cells were only transfected with constructs for Alk3CA. As in the case of the Tbx2 promoter, repression by the DNA binding-deficient mutant form of Tbx20 did not reach the level of the wild-type protein suggesting participation of direct transcriptional activities of Tbx20 wild-type protein in these assays. Wild-type Tbx20 protein acted as a DNA-dependent transcriptional activator on the Bmp/Smad-independent Nppa promoter (Habets et al. 2002) while the Tbx20-mutant protein did not show a transcriptional effect in this context (Fig. 6D). Hence, Tbx20 acts as a transcriptional activator of Bmp/Smad-independent promoters but as an inhibitor of Bmp/Smad-mediated transactivation.

DNA binding-independent inhibition of Bmp/Smad-mediated transactivation by Tbx20

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**Figure 7.** Tbx20 inhibits Bmp/Smad-signaling in a DNA-independent manner by binding to activating Smad1 and Smad5 proteins. (A) Pull-down of in vitro synthesed Flag-tagged Smad5, Myc-tagged Smad1 and HA-tagged Smad4, Smad6, and Smad7 proteins by GST fusion proteins bound to GSH-agarose beads and subsequent detection by anti-Flag, anti-Myc and anti-HA immunohistochemistry on Western blot. All GST fusion proteins harboring a T-domain retain Smad5 and Smad1 proteins on the column. (B) Co-immunoprecipitation of Myc-tagged Tbx20 and Flag-tagged Smad5 from HeLa cells co-transfected with expression constructs for the two proteins in presence and absence of an expression construct for constitutively active Bmp receptor Alk3 (Alk3CA). Arrow indicates the presence of Smad5 protein after IP with anti-Myc antibody and subsequent detection with anti-Flag antibody. The lower band in the Western blot represents the Ig heavy chain. (C) Co-immunoprecipitation of Flag-tagged Smad5, HA-tagged Smad4, and increasing amounts of Tbx20,HA in the presence of AlkCA from HeLa cells. Subsequent immunohistochemistry for different epitopes on Western blot detects competition of Smad4 binding to Smad5 by Tbx20. (D) Co-immunoprecipitation of Myc-tagged Smad1 and HA-tagged Smad4 from HeLa cells co-transfected with expression constructs for the two proteins, and subsequent addition of in vitro translated Tbx20 protein to Protein A bound complexes. Anti-HA immunohistochemistry on Western blot detects competition of Smad4 binding to Smad1 by Tbx20.
may rely on physical interaction and/or functional interference with the transcriptional activators Smad5 or Smad1. *In vitro* pull-down assays with GST-Tbx20 fusion proteins showed that Tbx20 directly binds to the activating Smad5 and the closely related Smad1, but not to the regulatory Smad4 and the inhibitory Smads, Smad6 and Smad7 (Fig. 7A, Suppl. Fig. 7A). Binding was mediated by the T-box of Tbx20 as shown by GST pull-down assays with fusion constructs of GST and various Tbx20 protein fragments (Fig. 7A). Tbx20 binding to Smad5 also occurred in a cellular context as shown by co-transfection/co-precipitation experiments in HeLa cells (Fig. 7B). Since Smad4 is a necessary co-factor for nuclear translocation and transcriptional activation by Smad1/Smad5 we investigated whether Tbx20 binding to Smad1/Smad5 competes with Smad1/5-Smad4 complex formation. We transfected expression constructs for Alk3CA, Smad4.HA, Smad5-Flag and increasing amounts of HA-tagged Tbx20 into HeLa cells and precipitated Smad5/Smad4/Tbx20 complexes with anti-Flag antibodies. Detection of HA-epitopes on Western Blots revealed a decrease of co-immunoprecipitated Smad4.HA protein in the presence of increasing amounts of Tbx20 (Fig. 7C).

In an alternative assay, we transfected expression constructs for Myc-tagged Smad1 and HA-tagged Smad4 into HeLa cells and precipitated Smad1/Smad4 complexes by anti-Myc antibodies. Addition of *in vitro* translated Tbx20 protein to resuspended immunocomplexes resulted in a complete release of Smad4 from the complex (Fig. 7D). Thus, Tbx20 effectively competes with Smad4 for binding to Smad1/Smad5 explaining the DNA-independent inhibition of Bmp/Smad-mediated activation of target promoters including *Tbx2*.

**Discussion**

Mammalian heart development is characterized by an intricate spatial and temporal interplay between morphogenetic programs controlling cellular proliferation and differentiation processes. Bmp-signaling and T-box factor transcriptional activities have been identified to govern some of these programs but the mechanisms that direct and restrict signaling activities and link them to transcriptional circuits to achieve compartmentalization of the heart tube have remained elusive. Our study suggests that the T-box transcription factors Tbx2/Tbx3 and Tbx20 are targets and regulators of Bmp/Smad-signaling, respectively. Tbx20 promotes cardiac chamber formation, and simultaneously suppresses Bmp/Smad-transcriptional activity in the rest of the heart tube in a novel DNA-binding independent manner, thereby restricting Tbx2 activity to the prospective AV canal region. In turn, Tbx2 together with Tbx3, locally stimulate development of the AV canal myocardium and cushions, possibly by up-regulating Bmp-signaling.
Tbx20 restricts Tbx2 to the prospective AV canal by attenuating Bmp/Smad-signaling

Expression analyses, embryological manipulation in the chick and genetic ablation experiments in the mouse have revealed a range of Bmp-dependent processes during amniote heart development (van Wijk et al. 2007). Bmp-signaling is broadly activated in the precardiac mesoderm and the cardiac crescent and is required for differentiation of progenitors of the first and second heart field into cardiomyocytes (Monteiro et al. 2004; Klaus et al. 2007). Ablation of Bmpr1a in early mesoderm resulted in a complete failure to establish cardiomyocytes expressing conserved core cardiogenic factors including Tbx5, Tbx20, Nkx2.5 and Gata family members, implicating them as downstream mediators of early Bmp-signaling. After cardiac specification, Bmp-signaling is redeployed for specification of the AV canal. Conditional deletion of Bmp2 showed that Bmp-signaling is required to establish an AV myocardium, to enhance formation of the cardiac jelly and to induce endocardial EMT (Ma et al. 2005). Bmp2 regulates expression of the transcriptional repressor Tbx2 in the AV canal: Tbx2 expression is lost in Bmp2 mutants, and Tbx2 is induced by Bmp2-loaded beads in chick epiblast cultures (Yamada et al. 2000; Ma et al. 2005). Our analysis of the regulatory region of the Tbx2 gene identified a genomic fragment that is sufficient to direct AV canal expression in vivo. This fragment is rich in Smad-binding sites and responsive to Bmp/Smad-signaling in vitro, strongly arguing that Tbx2 is a direct target gene of this pathway in the heart.

Analysis of the regulatory region of tbx2b in the zebrafish heart has identified Foxn4 and Tbx5 as activators of tbx2b expression (Chi et al. 2008). Deletion of (conserved) TBE and Foxn4 sites in the Tbx2 upstream region of the mouse showed that these sites are not relevant for cardiac Tbx2 expression, suggesting that AV canal restriction of Tbx2 might have been achieved by different molecular pathways in vertebrate evolution.

Expression of Tbx2 in the early heart tube prevents chamber formation (Christoffels et al. 2004b), illuminating the necessity to prevent premature activation of Tbx2 by the first wave of Bmp/Smad-signaling in the cardiac crescent, the heart tube and the prospective chambers. In Tbx20-mutant hearts, Tbx2 is induced in the developing cardiac crescent and throughout the linear heart tube (Singh et al. 2005), demonstrating that temporal and spatial restriction of Tbx2 to the developing AV canal is not achieved by positive regulatory inputs, but by Tbx20-mediated inhibition of broad activation in regions outside the AV canal. Absence of TBE sites from a minimal Tbx2 promotor fragment that is sufficient to recapitulate cardiac Tbx2 expression in wild-type and Tbx20-mutant embryos strongly argues against Tbx20 acting as a transcriptional repressor in this context, as indicated by ChIP experiments (Cai et al. 2005).

To our surprise, we discovered that Tbx20 binds to activating Smad1 and 5 and sequesters them from binding to the common Smad4, abolishing the formation of transcriptionally active Smad1/5,Smad4 complexes. Hence, transcriptional modulation of target gene expression by T-box transcription factors may not only rely on the presence of
T-box genes in cardiac regionalization

TBE sites that feature a particular orientation and spacing. Direct binding and sequestration of transcriptional regulators by the conserved T-domain suggests another level of complexity of transcriptional regulation and establishes the T-box as a versatile interface both for DNA and protein interaction.

Tbx20-mediated sequestration of activating Smads may be one of several mechanisms that synergize to shut off Bmp/Smad-signaling after cardiac specification of the lateral plate mesoderm. In Nkx2.5-mutant hearts, Bmp/Smad-signaling is dramatically augmented and expanded suggesting that Nkx2.5 represses Bmp/Smad-signaling (Prall et al. 2007). Smad6 that is expressed in the cardiac crescent stably binds to activated type I receptors and competes with regulatory Smad4 for receptor activation (Imamura et al. 1997). Moreover, similar and likely in addition to Tbx20, Smad6 specifically competes with Smad4 for binding to receptor-activated Smad1, yielding an inactive Smad1/Smad6 complex (Hata et al. 1998). Since Tbx20, Nkx2-5 and Smad6 are targets of Bmp-signaling in the cardiac crescent, they may be part of a concerted negative feed-back loop.

Since Tbx2 is known from other systems like the limb to respond to high levels of Bmp only (Suzuki et al. 2004; Behesti et al. 2006; Yang et al. 2006), all these mechanisms may collectively dampen Bmp/Smad-signaling to a level insufficient to activate Tbx2. Intriguingly, we noted that ectopic expression of Tbx2 and Tbx3 resulted in increased expression of Bmp2, Bmp4 and Smad-target genes (e.g. Id). Thus Tbx2/Tbx3 may activate a feed-forward loop of Bmp-signaling that further increases Tbx2/Tbx3 expression, and thereby development of the AV canal.

**Tbx20 and Tbx2/3 regulate chamber versus AV canal development**

Previous analyses by a number of research groups revealed the crucial role of *Tbx20* in cardiac chamber formation in vertebrates (Cai et al. 2005; Singh et al. 2005; Stennard et al. 2005; Takeuchi et al. 2005). *Tbx2* expression, normally restricted to the AV canal (Habets et al. 2002) was expanded into the entire heart tube of Tbx20-deficient embryos arguing that chambers have been lost at the expense of an AV canal. Since ectopic expression of Tbx2 in the pre-chamber heart tube resulted in a similar phenotypic outcome (Christoffels et al. 2004b), it was hypothesized that de-repression of Tbx2 fully explains the cardiac phenotype in Tbx20-deficient embryos. Our analysis of Tbx20/Tbx2 double mutant embryos revealed that loss of Tbx2 does not rescue the early cardiac arrest, identifying Tbx20 as a positive factor for chamber formation independent from Tbx2. However, given the dominant chamber differentiation-blocking capacity of Tbx2 (Christoffels et al. 2004b), the ectopic expression of Tbx2 in Tbx20 mutants is likely to add to its cardiac phenotype. This is further supported by biochemical studies that show that the transcriptional repressor Tbx2 can compete with transcriptional activators of the T-box family including Tbx5 and Tbx20 for binding to conserved DNA-elements such as TBE sites in the Nppa promoter (Habets et al. 2002). Thus, Tbx20 has a dual role in cardiac development, acting to stimulate formation of chambers and to confine repressor Tbx2 to the AV canal.
Tbx2 and Tbx3, a pair of evolutionary closely related T-box proteins, share identical biochemical properties (reviewed in (Naiche et al. 2005). Co-expression in the AV canal argues for a role of these genes in regionalization of the simple heart tube and establishment of the AV canal phenotype. In gain-of-function scenarios both Tbx2 and Tbx3 were able to prevent chamber differentiation (Christoffels et al. 2004b; Mommersteeg et al. 2007) and to suppress a broad spectrum of working myocardium associated genes, including sarcomere components and mitochondrial genes. Ectopic expression of Tbx3 in atrial myocardium led to induction of a set of genes associated with the AV canal / conduction system in the atrial working myocardium. Individual loss of function of either Tbx2 or Tbx3 did not have a major impact on AV canal development (Harrelson et al. 2004; Ribeiro et al. 2007; Bakker et al. 2008; Mesbah et al. 2008). However, Tbx2/3 double mutant embryos largely failed to establish a morphological AV canal indicating that the two genes act redundantly in this process, possibly augmented by other factors including Id genes that suppress chamber differentiation (Moskowitz et al. 2007). Together, these data indicate that Tbx2 and Tbx3 coordinately regulate the AV gene program by suppressing working myocardial and by activating AV-specific features.

Tbx2/Tbx3 double mutant embryos failed to establish AV swellings (cushions), which are the precursors of the valves, and contribute to the septal structures and to the fibrous insulation. The Tbx20 mutants formed cardiac jelly in the entire tube, whereas Tbx2/Tbx20 double mutants did not, suggesting that Tbx2 triggers cardiac jelly formation. These data also suggest that Tbx20 that has been implicated in later aspects of cushion development (Shelton and Yutzey 2007) is not required for the initial steps of cushion formation. Furthermore, we observed a striking mesenchymal layer in the atria of mice ectopically expressing Tbx3 in the atria. Together, these observations strongly suggest that Tbx2 and Tbx3 in the AV canal are required and individually sufficient to initiate cushion formation and EMT. The mechanism of cushion formation has been extensively studied, and important roles for ligands and receptors of the Tgfß-superfamily have been exposed (Bartram et al. 2001; Camenisch et al. 2002; Sugi et al. 2004; Mercado-Pimentel and Runyan 2007). Bmp2 expression in the AV canal myocardium is both required and sufficient to induce cushion formation (Sugi et al. 2004; Ma et al. 2005). Bmp2 was selectively upregulated in the Tbx3-expressing atrial myocardium, and is likely to be a pivotal downstream mediator of cushion induction by Tbx3. This is further confirmed by enhanced expression in the ectopic mesenchyme of Tgfß2 and other Bmp2 target genes that are required for cushion formation (Bartram et al. 2001; Camenisch et al. 2002; Sugi et al. 2004; Mercado-Pimentel and Runyan 2007). However, alternative possibilities could be considered, because Bmp2 is not absent from Tbx2/Tbx3 double mutant hearts that fail to initiate cushion formation. One possibility is that in addition to its role of directly signaling to the endocardium through receptor Bmpr1a, Bmp2 may act through Tbx2 to activate another intercellular signaling pathway, such as the Notch pathway also implicated in cushion formation (Timmerman et al. 2004).

Together, our loss- and gain-of function analyses suggest that Tbx2/Tbx3 and Tbx20 antagonistically regulate regionalization of the heart. Tbx20 promotes chamber formation
whereas Tbx2/Tbx3 directs the AV canal phenotype in the growing heart tube. Chamber and AV canal formation may be coupled and localized by the antagonistic regulation of Bmp/Smad-signaling pathway by these T-box transcription factors.

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References


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Supplementary data

Supplementary materials and methods

Histological analysis
Embryos were embedded in paraffin wax and sectioned to 10 µm. For histological analyses sections were stained with Hematoxylin and Eosin.

In situ hybridization analysis
In situ hybridization analyses on whole embryos and on paraffine sections using digoxigenin-labeled antisense riboprobes were performed according to standard protocols (Wilkinson and Nieto 1993; Moorman et al. 2001). Details of probes are available upon request. Stained whole mounts were transferred into 80% glycerol prior to documentation on a Leica M420 microscope with a Fujix digital camera HC-300Z. Sections of in situ hybridizations were photographed using a Leica DM5000 microscope with a Leica DFC300FX digital camera. All images were processed in Adobe Photoshop CS.

Microarray analysis
Left atria of six Cre4-CT mice and six control (Cre4) mice (male, 6 weeks) were dissected and snap frozen in liquid nitrogen. Total RNA was isolated and purified using single prep nucleospin columns according to the manufacturer’s instructions (Macherey-Nagel). RNA quality was checked using a bioanalyzer (Agilent Technologies). 250 ng of total RNA was used for biotin-16-UTP cRNA labeling and amplification using the Illumina RNA amplification kit (Ambion Inc., Austin, Txs). Labeled RNA was hybridized to Illumina MouseRef-6 BeadChip following the manufacturer’s instructions (Illumina Inc., San Diego, CA). The arrays were scanned using an Illumina Bead array reader confocal scanner. Beadstudio software was used to assess the individual array quality. Unprocessed intensity values were averaged per bead type, exported from beadstudio and subsequently normalized using VSN in R (Huber et al. 2002). Genes were tested for significant differential expression using the empirical Bayes moderated t-statistics test in the R-Limma package (Smyth 2004) at a 5% Benjamini-Hochberg false discovery rate (Reiner et al. 2003). We found that the expression of 737 transcripts was significantly reduced in atria of Cre4-CT mice, whereas 809 transcripts were significantly induced (threshold: P-value<0.05). A comprehensive analysis of this data set will be presented elsewhere.

Quantitative Real-Time PCR analysis
Quantitative real time PCR analysis was performed as described before (Hoogaars et al. 2007). In short, total RNA was isolated from left atrial appendices of 4 week old adult mice using the RNeasy Mini Kit according to the manufacturer’s protocol (Qiagen). cDNA was reverse transcribed from 300 ng total RNA using the Superscript II system (Invitrogen) and expression of different genes was assayed with quantitative real time-PCR using the Roche 480 Lightcycler. Relative start concentration \( N(0) \) was calculated using the following equation: \( N(0)=10^{log(threshold)-Ct \ (mean \ Eff)} \). Values were normalized to Gapdh expression levels. Primers sequences can be provided upon request.
**Mutagenesis**

For construction of a DNA-binding deficient mutant form of Tbx20 conserved amino acid residues in the DNA-binding region were mutated based on reports for the Tbx2 protein (Habets et al. 2002; Lingbeek et al. 2002). Primer sets used for making the DNA binding dead mutant were:

- ATCACCAAGGTCTGGCAGAGGATGTTCCCCACCACATCC converting arginine 126 to glutamic acid,
- ATCACCAAGGTCTGGCAGGGAGATGTTCCCCACCACATCC converting arginine 127 to glutamic acid and
- ATCACCAAGGTCTGGCAGGGAGATGTTCCCCACCACATCC converting arginines 126 and 127 to glutamic acid residues. pcDNA3.1Tbx20.HA was used as a template for the amplification. Either of the three primers was used for the amplification of the entire plasmid using PfuTurbo DNA polymerase (Stratagene QuickChange XL Site-Directed kit manual). Positive clones were sequenced to confirm the mutation, and protein synthesis was analyzed in transfected Hela cells to check efficiency of expression. Details on all other constructs upon request.

**In vitro transcription/in vitro translation**

Coding regions of mouse Tbx20, Smad4, Smad6 and Smad7 were amplified by PCR from the respective cDNAs and inserted in pSP64 modified to contain 5'-ß-globin leader and 3'-ß-globin trailer sequences as C-terminal fusion proteins with Myc or HA-epitope tag (Kispert and Herrmann 1993). SP6-coupled in vitro transcription/translation kit (TNT, Promega) was used for synthesis of the proteins in wheat germ lysate.

**Expression in cell lines**

For cytomegalovirus promoter/enhancer-driven expression of Tbx20, Smad1 and Smad5 proteins in cells, the globin leader/cDNA/globin trailer cassette of pSP64 was shuttled into HindIII and EcoRI sites of pcDNA3 (Invitrogen). Constructs were transfected in HeLa cells using the calcium phosphate method and in NIH3T3 cells employing Fugene reagent (Roche).

**In vitro reporter assays**

Luciferase reporter assays were used to determine transactivation properties of Tbx20 on various promoter fragments. Promotor fragments cloned in pGL2 or pGL3-luciferase constructs (Promega) were the 6 kbp genomic Tbx2 fragment and deletions thereof (pGL3basic.Tbx2-Luc), a minimal Bmp-responsive Msx2 promoter element (pGL2basic.Msx2-Luc) (Brugger et al. 2004) a short 91 bp Bmp-response element derived from the Id1 promoter (pGL3.BRE2-Luc) (Monteiro et al. 2004) and a 0.7 kbp Nppa genomic fragment (pGL3basic.Nppa(0.7)-Luc) (Habets et al. 2002). Reporter constructs were co-transfected with expression constructs for HA-tagged mouse Tbx20 (pcDNA3.Tbx20.HA), HA-tagged mouse DNA-binding-deficient form of Tbx20 (pcDNA3.Tbx20.HAmut) alone or in the presence of Flag-tagged mouse Smad5 (pcDNA3.Smad5.Flag) and constitutively active Bmp-receptor Alk3 (Alk3CA) (pCS2.BmpR1a.CA)(Wessely et al. 2001). Constructs were transfected into NIH3T3 cells (6×10^5 cells per well of 6-well plates) with the Fugene HD
transfection reagent (Roche). 40 ng of pCMV.ßGal vector were co-transfected to normalize the transfection efficiency by colorimetric determination of X-Gal turnover in the ß-Gal assay. After 48 h of further culture, cell lysates were prepared and the luciferase and ß-galactosidase activities were measured. All transfections were performed in duplicates and experiments were repeated at least three times. After normalization, the mean luciferase activities and standard deviations were plotted as “fold activation” when compared with the empty expression plasmid.

**Immunoprecipitation**

To determine binding of Tbx20 to Smad5 by co-immunoprecipitation experiments, Hela cells were either transfected with expression constructs for HA-tagged mouse Tbx20 (pcDNA3.Tbx20.HA) alone or in the presence of Flag-tagged mouse Smad5 (pcDNA3.Smad5.Flag) and constitutively active Bmp-receptor Alk3 (Alk3CA) (pCS2.BmpR1a.CA) (Wessely et al. 2001). Transfections were performed using the calcium phosphate method in 10-cm dishes at 50-60% confluency. After 48 h, cells were lysed in 1000 µl of Nonidet P-40 buffer, cellular debris was precipitated by centrifugation for 20 min at 4°C. The supernatant was split for immunoprecipitation with anti-HA and anti-Flag antibody (2.5 µg). After 1 h of incubation, 25 µl of protein G sepharose beads (Amersham Biosciences) were added for 2 h at 4 °C followed by the precipitation and washing of beads. Beads were boiled in SDS-loading buffer and eluted proteins analyzed by Western blot.

To determine competition of Tbx20 and Smad4 for binding to Smad1 in co-immunoprecipitation experiments, Hela cells were transfected with expression constructs for HA-tagged mouse Smad4 (pcDNA3.Smad4.HA) and myc-tagged mouse Smad1 (pcDNA3.Smad1.Myc). Transfections were performed by the calcium phosphate method in 10-cm dishes at 30% confluency. After 48 h, cells were lysed in 500 µl of Nonidet P-40 buffer, cellular debris was precipitated by centrifugation for 15 min at 4°C. The supernatant was split in three aliquots for immunoprecipitation with anti-Myc antibody (2.5 µg). After 2 h of incubation, 30 µl of proteinA agarose beads (Amersham Biosciences) were added for 1 h at 4°C followed by the precipitation and washing of beads. Beads were resuspended in 300 µl Nonidet P-40 buffer, and 0, 10 and 50 µl of in vitro translated Tbx20 protein was added. After 2 h incubation at 4°C, beads were precipitated and washed. Proteins were released by boiling in SDS-buffer, and separated by SDS-PAGE. After Western blotting, Myc-tagged proteins (Smad1.Myc, Tbx20.Myc) were detected by anti-Myc, while Smad4.HA was detected by anti HA-immunohistochemistry.

To determine competition of Tbx20 and Smad4 for binding to Smad5 in co-immunoprecipitation experiments, HeLa cells were transfected with expression constructs for Alk3CA, HA-tagged mouse Smad4 (pcDNA3.Smad4.HA), Myc-tagged mouse Smad5 (pcDNA3.Smad5.Myc) using the calcium phosphate method as described. Specific proteins were detected with immunohistochemistry on Western blots as described.

**GST pull down**

For GST pull-down experiments, GST or GST fusion proteins with five different deletion domains of Tbx20 were expressed in E. coli strain BL21 and bound in the presence of DNaseI to glutathione sepharose 4B beads (Amersham Biosciences) as described (Leger et al. 1995). An aliquot of the washed and equilibrated beads, now carrying GST or GST fusion protein, was incubated with one-tenth of an extract of HeLa cells transfected with pcDNA3.1 expression constructs for Smad5.Flag or Smad1.Myc protein (from 10cm plates), or with an
aliquot of in vitro translated protein of Smad4.HA, Smad6.HA and Smad7.HA in interaction buffer (20 mM HEPES, pH 7.9, 100 mM NaCl, 10 mM KCl, 5mM MgCl$_2$, 0.5 mM EDTA, 5% glycerol, 0.05% Triton X-100, and 1 mM DTT). After extensive washing, the proteins were eluted and analyzed by SDS-PAGE and Western blot.

**Electrophoretic mobility shift assay**

EMSA was done as previously described (Farin et al. 2007). The probe for Tbx20 binding was generated by annealing the two oligonucleotides BS.dirF, 5'-GATCCGGAGGTGTAAGGTGAAAGGA-3'; and BS.dirR, 5'-GATCTCCCTTTCCACACCTTCACACCTCCG-3'. Protein for the binding assay was prepared using TNT SP6 High-Yield protein expression system (Promega) (Farin et al. 2007).

**In vivo reporter assays**

The -6mTbx2-Eyfp construct for generating transgenic Tbx2 promoter-reporter lines (Fig. 5A) was generated by inserting a 6 kbp Tbx2-promoter fragment (from -5.557 bp to +310 bp relative to the human TBX2 transcription start site) into expression vector pCS2, upstream of Eyfp (Venus) (Nagai et al. 2002), removing the CMV promoter. The -3.6mTbx2-Eyfp and -2.7mTbx2-Eyfp constructs were generated by truncation of the -6mTbx2-Eyfp construct using the restriction sites NheI and AflIII, respectively (Fig. 5A). To generate the -1.5mTbx2-Eyfp and -1.3mTbx2-Eyfp constructs, the -3.6mTbx2-Eyfp construct was restricted with AflIII and BglII, after which a fragment was cloned in between with an artificial AflII site at -314 bp and -63 bp, respectively. Within the -1.5mTbx2-Eyfp construct, a conserved LEF1/Tgf binding site CTTTGTT (Arce et al. 2006) at -2620 bp was mutated into CccgcGcgGT to generate the -1.5(LEFmut)mTbx2-Eyfp construct. Vector sequences were removed and constructs were injected into pronuclei of zygotes of FVB mice.
Supplementary figures

Supplementary Figure 1. Analysis of AVC formation in an allelic series of Tbx2 and Tbx3 embryos by cardiac morphology, histology and expression of the chamber marker Nppa at E9.5; genotypes are as indicated. (A-F) Left lateral views of whole embryos. Boxes mark the heart regions to be magnified in the following figures (G-L). (M-R) Histological analysis of sagittal sections through the left atrium (la), atrioventricular canal (avc) and left ventricle (lv) by hematoxylin and eosin staining (H&E). (S-D') in situ hybridization analysis of Nppa expression in whole hearts in dorsal views (S-X) and in left lateral views (Y-D'). Embryos with a loss of three functional alleles of Tbx2 and Tbx3 exhibit expansion of chamber myocardium in the AVC, and lack of AVC cushion formation, while single mutants appear phenotypically normal in this respect.
Supplementary Figure 2. Tbx3 induces expression of genes involved in cardiac cushion formation and valve development. Serial sections of E17.5 hearts of CT and Cre4;CT mice, showing induced myocardial expression of transgenic TBX3, reduced myocardial expression of Smpx and Bmp10, induced myocardial expression of Cacna2d2 and Hcn4, and induced mesenchymal expression of Odd1, Id3, Meox1 (also myocardial), Frodo (also myocardial), Bmp6, Smad6 and Collagen III. ra, right atrium; san, sinoatrial node. Black bar, 100 µm.
Supplementary Figure 3. Cross-sections of E12.5 mouse hearts showing expression of genes up-regulated in Cre4;CT atria (micro-array) in the developing AVC and AV cushions and valves. In situ hybridization shows expression of Tbx3 in AVC myocardium (green arrow heads) and endocardial cushions (asterisk). Aldh1b1 expression is absent from the AVC, whereas Hnt, Cacna2d2 and Hcn4 are expressed in the AVC and sinus node. We observed broad expression of Sox9, Mdk, Fbln2, Txndc5, Lum and Lef1 in the AV cushions (black asterisk) and the valve leaflets, whereas expression of Meox1, Id3, Nkd2, Fgfr2 and Frzb was more restricted.
Supplementary Figure 4. Sequence analysis of the 6 kbp promoter region of Tbx2 identifies conserved elements.

Reference sequence NCBI July 2007: >ref|NT_096135.5|Mm11_95772_37:51144025-51167297 Mus musculus chromosome 11 genomic contig, strain C57BL/6J.

Positions of restriction sites and putative binding sites refer to the middle of a site.

-6mTbx2 promoter

The first nucleotide (G) and last (A) are depicted in green. Total length is 5867bp, from -5557 to +310 relative to the transcription start site.

5'-GGGATGTTAGAAGCTTCTGATAGCATGACCCCTGCACCCACCCTTCTGATACCAAGAGTATGGGACCTTTCTCAGCCCCGCAGGGGACCTACGGACCCTGGCCCAGCCCTTTACAGTGGGCAAAAAAACAGATGTGGCCTCCTCCAGTCAGGGAAAGCAGTGGGTGTAGCGCTGGACGTGCTGGCTGAGCTTGCAGGAC

CAGAC

GACCTCGCAGAGCAGACTGAGAAGGAGCATTTGAAATTTCAGTGATCGGTTCTAAATGTTAAAGATCGGAGACTAACCCCCACTCATTCCACCCTCCTTTTATTGATGACAAGCCTGAGCCCCCAAGAAGGGCACTGGCCTACACTCCGAGCTGCACCTCCTGGGACCAAGGGCTCAGGGTAGCAAGTATTTAGCAGTTTTTTAGGAGTTATACGATTCCTAAAGCTCCCCCAAATCGACTATAACCTATCTTCATACCGCGTGATCTCTCACTATGCAAGAAATGCCCGCCCACCGGAGCAAGTGTGTCCCTGACAGAGACAGTTTGGTTGGGCCTCGCCTTTGCATTAACCTGGAAGGGCCCGATTATCTGTGATAGCGGCGGACTGCGCAGCAAGGCGCCGCTCGAGAGGCTGCTTAGGCCCCCCAGCACCAGTTTTCGATGCTACAAGGATTCTAAGAAAATTTTGTTGGGGATGGGTCGATAAAGAATCTTAAGAGCGGCTAAAGCAGGTCATCCAGGTGACTACGTGATGGGGGTGGGGGGGTGGCTGGAGCTGTTGGTGGAAACCCAACCTGCCCTGAAAACGTCTCTGACCAGTCGCAGGTCCTTAGCTCTATAGGTGGGATTCAGATCCCCTAATGTAGCCACTGGGC
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GACAGCTGCGGATTTCGTGGACCAATAACCTTCAAGCTGCTTCTCGCAGAAAGCTGAGACTGACTACTCC
CTCTGTTTACCTCCTCATACCTTCTCCTTTTACAGGACATACCTCGTTCTGCAGAAACGCTGACATAC
TCAAATTGGGACTGGCAAGCTGAGACTGACTACTCCCTCCCTTCTCTTACAGGACATACCTCGTTCTGC
AGAAACGCTGACATAC

TGAATTGGAAGCAGTGCAAAGCTGCAATGGATCTCTCACCGGCTGCGAACACCGCGAAGCTAGAGCTGAA
GGTTCACTGTGGGCAGAGAAAAGAGTGAGAAAAGCGACCTAGGAGTAAGGTCCTGAAGAAAAACCCAGTG
GGCAGGCGCCACAGGGCTTTCTGCTTCTCAAGTGGGACACAGAGGAGGAGGGAGATGCACTGTTTCCACT
GCTCCAAAGGGCTTATTCTCCCTTCTCTTACAGGACATACCTCGTTCTGCAGAAACGCTGACATAC
TCAAATTGGGACTGGCAAGCTGAGACTGACTACTCCCTCCCTTCTCTTACAGGACATACCTCGTTCTGC
AGAAACGCTGACATAC

-314
CTTTCCAGTGCCCTCTGAGTTCAATGCTTTGCACTTGGGGCCGGCATGGCGCTAGGGGGTCCTTCCCCAAG
GCCCCGCCCCCCGGCCCCGGCTCCCAGGAGCCTGGGCCGGATGTCCCGATGAGAGAGCCGGCGCTGGCAGCC
A

-3'

Transcription start site +1 = CAGAGATCA, conserved across species. Translation start site at +316 (ATG), conserved across species. The 5' untranslated region has been underlined. Transcription and translation start site of mouse have been inaccurately assigned in the NCBI database.

0.9 kb NheI-AflII enhancer
This fragment of 929 bp reaches from the unique restriction site NheI at -3233 to the unique restriction site AflII at -2305 bp. Its sequence has been marked in light grey.

-1.5 kb construct: promoter starts at -314
It contains the 0.9 kb NheI-AflII enhancer, which has been cloned upstream of a -0.6 kb promoter fragment from -314 to +310. The 5' nucleotide of the proximal promoter has been marked black.

-1.3 kb construct: promoter starts at -63
It contains the 0.9 kb NheI-AflII enhancer, which has been cloned upstream of a -0.4 kb proximal promoter fragment from -63 to +310. The 5' nucleotide of the proximal promoter has been marked light grey.

Putative binding sites previous literature on Tbx2 promoter:
Tbx5 BE at -173: AAGGTGTCGGAA (anti-sense)
Foxn binding site at -84: TTTACGCTTT (anti-sense)

**Novel localized putative LEF1 and T-box binding sites present study**

1) LEF1 binding site at -2620: 5'\-GCTTTGTT\-3' (reviewed in Arce L 2006. Oncogene 25:7492-7504), found to be conserved between human and mouse using Transfac matrices. To inactivate the binding site in the context of the -1.5(LEFmut)mTbx2-Eyfp construct it has been mutated into: GCcgcGcg (Giese K 1992. Cell 69:185-195).

2) Non-consensus putative Tbx5 site (for consensus sites see Sinha S 2000. Gene 258:15-29; Farin HF 2007. J Biol Chem 282:25748-59), localized using rVISTA and Transfac matrices, at -2615: 5'\-TGTTGTGGA\-3'. It partially overlaps with the LEF1 binding site. Within the -1.5(LEFmut)mTbx2-Eyfp construct the site has been inactivated by mutation to: 5'-cGcgGTGGA-3'.

**Novel localized putative SMAD binding sites present study**

**Consensus Smad Binding Elements (SBEs)**

5'-GTCTG-3' or 5'-CAGAC-3' was the original consensus binding sequence; 5'-TGCTG-3' or 5'-CAGACA-3' was indicated by Jonk LJC 1998. J Biol Chem 273:21145-52. However, also 5'-GTCT-3' or 5'-AGAC-3' has been published to be sufficient (Shi Y 1998. Cell 94:585-594; Zawel L 1998. Mol Cell 1:611-617; reviewed in Massagué J 2000. EMBO J 19:1745-1754). All sites have been indicated in the regulatory sequence.

**GC-rich palindromes shown to be important for SMAD activation**

5'-GGCGCC-3' (Korchynskyi O and ten Dijke P 2002. J Biol Chem 277:4883-4891)

**Non-consensus SMAD-binding sequences**

5'-GTGGAGCCTG6CGCCGGCTG6CCGA6CGC-3': Sequence at -2608, found to contain several SMAD binding elements which are conserved between mouse-human-chicken, using rVISTA and Transfac matrices. The sequence has been underlined.
Supplementary Figure 5. Identification of cardiac regulatory elements by deletion analysis of the Tbx2 promoter in transgenic embryos. RankVISTA alignment of human against mouse Tbx2 regulatory sequences show conserved regions. Conserved T-box binding elements (TBE), LEF and Foxn sites are marked by boxes. Top panels show pictures of the Tbx2 expression pattern in E10.5 hearts visualized by whole mount in situ hybridization. Below these are fluorescence pictures of hearts of representative embryos with constructs depicted in the right panel. White arrowheads depict the atrioventricular canal (AVC), grey arrowheads the outflow tract (OFT). Grey lines indicate Tbx2 regulatory sequences, boxes represent exons, green boxes reporter gene Egfp. la, left atrium; lv, left ventricle; ra, right atrium; rv, right ventricle.
Supplementary Figure 6. Identification of regions of the Tbx2 promoter conferring transcriptional activation by Bmp/Smad-signaling and repression by Tbx20 in transactivation experiments in vitro by deletion analysis. Plasmids encoding constitutively active Bmp receptor Alk3 (Alk3CA), full length Smad5 protein, full length Tbx20 protein and the luciferase reporter constructs were co-transfected in NIH3T3 cells and luciferase activity determined and normalized as fold activation over the reporter alone. Numbers indicate µg of plasmids for the reporter -6.0Tbx2-TBE-luc (A), -5.0mTbx2-luc (B), -3.6mTbx2-luc (C), -2.7mTbx2-luc (D) and -6.0mTbx2-0.9-luc (E), and the expression plasmids for Alk3CA, Smad5 and Tbx20 co-transfected into NIH3T3 cells.
Supplementary Figure 7. Biochemical analyses of Tbx20 proteins. (A) Coomassie staining of Tbx20-GST fusion proteins shows that all proteins are expressed in and purified equally well from bacteria. N refers to the N-terminal protein domain, T to the T-box region and C to the C-terminal protein domain. (B) Western blot analysis of the HA-tag of different Tbx20 proteins in transfected HeLa cells shows that the wild-type (wt), and the non-DNA binding Tbx20mut1 (L126R), Tbx20mut2 (L127R) and Tbx20mut (LL126,127RR) forms are expressed equally well. (C) Electrophoretic mobility shift assay of wild-type and mutant Tbx20 proteins confirms that amino acid changes L126R and L127R abolish binding of the Tbx20 protein to the DNA probe. Note the absence of the supershift band (arrow) in the mutant situations (Tbx20.HAmut1, mut2, mut).
References supplementary data


Gene expression profiling of the forming atrioventricular node using a novel Tbx3-based node-specific transgenic reporter

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Abstract

The atrioventricular (AV) node is a recurrent source of potentially life-threatening arrhythmias. Nevertheless, limited data are available on its developmental control or molecular phenotype. We used a novel AV node-specific reporter mouse to gain insight into the gene programs determining the formation and phenotype of the AV node. In this reporter, green fluorescent protein (GFP) expression was driven by 160 kbp of Tbx3 and flanking sequences. GFP was selectively active in the AV canal of embryos, and AV node of adults, whereas all other Tbx3+ conduction system components, including the AV bundle, were devoid of GFP. These findings demonstrate that distinct regulatory sequences and pathways control expression in the components of the conduction system. Fluorescent AV nodal (Tbx3BAC-Egfp) and complementary working myocardial (NppaBAC336-Egfp) cell populations of E10.5 embryos and E17.5 fetuses were purified using fluorescence-activated cell sorting, and their expression profiles were assessed by microarray analysis. We constructed a comprehensive list of sodium, calcium, and potassium channels specific for nodal or working myocardium. Furthermore, the data revealed that the AV node and the working myocardium phenotypes diverge during development, but that the functional gene classes characterizing both subtypes are maintained. One of the repertoires identified in the AV node-specific gene profiles consists of multiple neurotrophic factors and semaphorins, not yet appreciated to play a role in nodal development, revealing shared characteristics between nodal and nervous system development. These data present the first genome-wide transcription profiles of the AV node during development, providing valuable information concerning its molecular identity.
Chapter 4

Introduction

The electrical impulse is initiated in the sinus node, propagated rapidly through the muscular tissues of the atria, delayed in the AV node, and then rapidly propagated through the ventricular conduction system, activating the ventricles from apex to base. The slow conduction through the AV node ensures sequential, synchronized contractions of the atria and ventricles (reviewed in Meijler and Janse, 1988; Anderson et al., 1981). This guarantees the ventricles to benefit from the atrial contraction, adding 15% to cardiac output, and prevents one to one conduction of potentially life-threatening supraventricular arrhythmias. Furthermore, the AV node is the major subsidiary pacemaker in case of failure of the sinus node (reviewed in Meijler and Janse, 1988; Mangoni and Nargeot, 2008). However, the AV node is also a recurrent source of tachy- and brady- arrhythmias, notably AV node reentrant tachycardia and AV block, requiring lifelong medication, ablation, or electronic pacemaker implantation. Knowledge regarding development (Virágh and Challice, 1977b; Moorman and Christoffels, 2003; Gourdie et al., 2003; Myers and Fishman, 2003; Stroud et al., 2007), anatomy (Anderson et al., 1981), and molecular background of the physiological function (Marionneau et al., 2005; Li et al., 2008; Boyett et al., 1996; Schram et al., 2002; Nerbonne and Kass, 2005) of the AV node has increased gradually. Nevertheless, insights into the transcriptional pathways regulating its development, and the gene expression profiles underlying its phenotype and function are limited.

The T-box transcription factor Tbx3 belongs to an evolutionary conserved family of factors, plays key roles in development and cancer, and is mutated in the ulnar mammary syndrome of congenital defects (Naiche et al., 2005). Tbx3 is specifically expressed in the components of the cardiac conduction system, i.e. the sinus node, AV node, AV bundle and bundle branches, throughout development and in the adult (Hoogaars et al., 2004), where it plays important roles in their formation (Hoogaars et al., 2007; Bakker et al., 2008). We analyzed the function of regulatory sequences of Tbx3 to gain insight into transcriptional regulation during conduction system formation. We identified a large regulatory fragment driving GFP reporter gene expression specifically in the AV node. Subsequently, this transgenic mouse model was used to study the transcription profiles of the developing AV node through microarray analysis, providing new insights into the molecular pathways underlying differentiation and function of the AV canal and maturing AV node.

Materials and methods

Transgenic mice

The transgenic working myocardium-specific promoter-reporter line NppaBAC336-Egfp has been described previously (Horsthuis et al., 2008). Bacterial artificial chromosome (BAC)
Development and phenotype of the AV node

clone 366H17, containing genomic sequences from -83 kbp to +77 kbp relative to the transcription start site of Tbx3, was obtained from a C57BL/6J mouse BAC/PAC library (CHORI, Oakland, CA). To generate the Tbx3BAC-Egfp construct, the sequence ATG.agc.ctc.t was replaced by GFP using the BAC modification method kindly provided by Gong and Heintz (Gong et al., 2002). An expanded Materials and Methods section is available in the online data supplement and at the end of this chapter.

Figure 1. Overview of Tbx3 regulatory sequences containing reporter constructs used in this study. (A) RankVISTA alignment of human against mouse Tbx3 genomic region covered by Tbx3BAC336H17. Bars depict evolutionary conserved segments where the heights scale with statistical significance [-log10(P-value)]. (B) After SstII restriction of the original Tbx3BAC and the modified Tbx3BAC-Egfp construct, an ethidium bromide stained gel (left) and Southern blot (right) depict the modified fragment. (C) Constructs used to generate transgenic mice. Fraction of founders (transg) expressing GFP (expr) and fraction of expressing transgenes expressing GFP in the heart (cardiac) and ectopically (ectopic), have been indicated. (D, H) Whole mount in situ hybridization of Tbx3. (E-F and I-J) Whole mount β-galactosidase staining of two independent founders carrying the -6/+2mLacZ construct. (G,K) Fluorescence pictures of mice carrying Tbx3BAC-Egfp construct. Black lines in A and C indicate Tbx3 regulatory sequences, boxes represent exons, dark blue or green boxes the reporter genes. CNCS, conserved non-coding sequence; M, λ-BsteII marker; UTR, untranslated region; lv, left ventricle; fl, forelimb; ra, right atrium; la, left atrium; rv, right ventricle; lsh, left sinus horn; rsh, right sinus horn.
Results

*Tbx3 activity is driven by distal regulatory sequences*

To assess the spatiotemporal activity of the proximal regulatory sequences of *Tbx3*, transgenic mice were generated in which a fragment ranging from 6 kbp upstream to 2 kbp downstream of the *Tbx3* translation start site drives *LacZ* reporter gene expression (Figure 1C). 5 of 6 independently obtained lines expressed *LacZ* in a pattern unlike that of *Tbx3*. In 4 of these lines *LacZ* was expressed mainly ectopically in the caudal part of the embryo and the forelimbs, whereas no expression was detected in the *Tbx3* tissues such as the mammary glands or heart (Figure 1D-F,H-J). These observations indicate that the regulatory sequences driving correct spatiotemporal *Tbx3* expression are located outside the 8 kbp regulatory DNA region.

![Figure 2. Comparison of expression patterns of endogenous Tbx3 and the Tbx3BAC-Egfp transgene.](image)

(A) Whole mount and (D-L) section in situ hybridization, and (B-C) fluorescence pictures. Arrowheads point at ulnar site limbs. hc, hypoglossal cord; hl, hindlimb; mg, mammary gland; mr, mandibular and maxillary region; wh, whisker; m, mandibula; san, sinoatrial (sinus) node; avb atrioventricular bundle; br, bronchi; aavc, anterior atrioventricular canal; lavj, left atrioventricular junction. See legend to Figure 1 for other abbreviations.
We next used a BAC clone containing Tbx3 flanked by 83 kbp upstream sequences and 77 kbp downstream sequences (Figure 1A-C). An enhanced green fluorescent protein (GFP) encoding reporter gene was inserted at the translation start site of Tbx3, thereby functionally inactivating Tbx3. 4 independent mouse lines carrying Tbx3BAC-Egfp showed comparable spatiotemporal GFP patterns (Figure 1G,K and 2B,C) highly similar to the pattern of Tbx3 (1D,H and 2A). In situ hybridizations on serial sections confirmed that in the eye, snout, mammary glands and limbs Tbx3BAC-Egfp mimicked Tbx3 expression (Figure 1G and 2A-I, Supplementary Table 1). However, the construct appeared inactive in the ear (Figure 1G and 2B,C) and the lung mesenchyme (Figure 2K,L), showing that not all DNA sequences regulating Tbx3 expression were present in the 160 kbp BAC.

**Tbx3BAC drives AV node-specific expression**

Tbx3 is expressed in the sinus node, AV node, AV bundle and bundle branches (Hoogaars et al., 2004), but Tbx3BAC-Egfp was not active in the Tbx3+/Cx43− sinus node, AV bundle and bundle branches (Figure 2J-L). In contrast, Tbx3BAC-Egfp was active in the AV canal from E9 onwards. Until E14.5 Tbx3BAC-Egfp was active in the dorsal, ventral and right side of the AV canal, colocalizing with the developing AV node (Virágh and Challice, 1977b), right atrioventricular ring bundle (Moorman et al., 1998), retro-aortic root branch (Moorman et al., 1998), and the anterior node-like structure (Blom et al., 1999) or retroaortic node (Anderson et al., 2008), respectively (Figure 2K,L and 3A-D). No activity was detected in the Tbx3+ left AV canal. Also the Tbx3+ AV mesenchymal cushions were devoid of GFP (Figure 4A). At E17.5, the area of activity in the AV junction (Figure 3E) further diminished, resulting in Tbx3BAC-Egfp activity progressively restricted to the AV node. In the adult, we detected GFP expression strictly limited to the Tbx3+/Hcn4+ tissue of the AV node and the anterior node, complementary to the Cx40+ AV bundle (Figure 3G-O).

To further verify AV nodal specificity of Tbx3BAC-Egfp, we performed patch-clamp experiments on single GFP+ and GFP− cardiac cells of E17.5 embryos. All action potential parameters, except overshoot, differed significantly between both groups (Figure 3P,Q and Supplementary Table 2). Importantly, all GFP+ cells (n=4) showed diastolic depolarization resulting in spontaneous activity, and exhibit the hyperpolarization-activated current (I_f), whereas all GFP− cells (n=5) had a stable resting membrane potential, were quiescent and lacked I_f (Figure 3P,Q and Supplementary Table 2). In conclusion, GFP+ cells had electrical properties typical for nodal cells and GFP− cells for working myocardial cells. The combined data of the marker expression analysis and patch-clamp experiments indicate that the 160 kbp Tbx3BAC-Egfp construct contains regulatory sequences driving specific AV nodal expression, but lacks additional enhancers driving expression in the sinus node, AV bundle and bundle branches.
Figure 3. Tbx3BAC-Egfp expression in the AV canal is progressively restricted to the developing AV node. (A-F) Atrioventricular fluorescence of mice carrying Tbx3BAC-Egfp shown from E9.5 till 3 weeks after birth. (G-O) In situ hybridization on serial sections confirmed that GFP was expressed only in the Tbx3⁺/Hcn4⁺/Cx40⁻ AV node, and not in the Tbx3⁺/Hcn4⁺/Cx40⁺ AV bundle. (P) Typical examples of action potentials (APs) from an E17.5 GFP⁺ and an GFP⁻ cell obtained by patch-clamp experiments. APs recorded from all GFP⁺ cells were spontaneous (n=4); APs from GFP⁻ cells were elicited by current injection through the pipette (n=5). Note the differences in time scale. Inset shows the maximal dV/dt, a measure for sodium current availability (Q) Membrane currents activated by 500 ms hyperpolarizing voltage-clamp steps from -40 mV (inset). Note the presence of the hyperpolarizing current (I_f) in the GFP⁺ cells (arrow). This current was never detected in non-fluorescent cells. avn(-pr), atrioventricular node(-primordium); oft, outflow tract; thd, thyroid gland; ravrb, right atrioventricular ring bundle; mv, mitral valve; tv, tricuspid valve; av, aortic valve; scv, superior caval vein; icv, inferior caval vein; vs, ventricular septum; bb, bundle branches. See legends to Figure 1 and 2 for other abbreviations.
Gene expression profiles of the developing AV node and working myocardium

The property of Tbx3BAC-Egfp to drive GFP expression specifically in the AV node myocardium allowed assessing the genome wide expression profiles of these cells during development. Nppa, the gene encoding atrial natriuretic factor, is specifically expressed in developing working myocardium complementary to the Tbx3+ conduction system (Moorman and Christoffels, 2003). The expression of GFP in the NppaBAC336-Egfp transgene recapitulates the Nppa expression pattern precisely (Horsthuis et al., 2008). Embryonic E10.5 Tbx3BAC-Egfp cells (E10.5 AV canal) were compared with age-matched pooled atrial and ventricular cardiomyocytes expressing NppaBAC336-Egfp (E10.5 WM (working myocardium)). Fetal E17.5 Tbx3BAC-Egfp (E17.5 AV nodal) cells were compared with E17.5 NppaBAC336-Egfp atrial working myocardium (E17.5 WM) (Figure 4A and 5A). For every sample, transgenic hearts were dissected, the GFP-positive tissues pooled, dissociated to obtain single cells, and purified using FACS (Figure 4B). Fluorescent cells comprised, on average, 1-4% of live-gated AV nodal, and 3-8% of live-gated WM cell populations, whereas

Figure 4. Purification of working myocardium- and AV node myocardium-specific myocyte populations for transcriptome analyses. (A) In situ hybridization and immunofluorescent labeling on sections of mice carrying the NppaBAC336-Egfp or Tbx3BAC-Egfp construct, respectively. (B) Successive steps of fluorescent cell preparation from dissection of the Egfp+ hearts till hybridization of amplified and labeled aRNA to the microarray. (C) FACS profiles of hearts of an E17.5 Tbx3BAC-Egfp+ transgene (right) and a wild type littermate (left). The percentage of Egfp+ cells in the live gate is shown. Arrowheads in A point at atrioventricular canal. See legends to Figure 1 and 3 for abbreviations.
Figure 5. Microarray experiments comparing E10.5 and E17.5 AV nodal myocardium with stage-matched working myocardium cell populations. (A) Scheme of microarray experiments. In present study experiment I and II were performed, after which data regarding III were deduced (For details about III, see Figure 6C). (B) Pie charts summarizing microarray experiment I (E10.5) and II (E17.5), showing number of transcripts enriched in AV nodal tissue relative to age-matched WM (N>W), transcripts enriched in WM relative to age-matched AV nodal myocardium (W>N), transcripts expressed above threshold (expr>thresh) in nodal myocardium and/or WM which are not differentially expressed between both groups, and transcripts not detected above threshold (expr<thresh). (C) Genes previously established to be differentially expressed between AV nodal and age-matched chamber/working myocardium at E10.5 and E17.5 were picked up in the array experiment.

post-sort analysis revealed 85-90% purity of GFP⁺ cell populations after flow cytometry (Figure 4B,C). Because the number of E10.5 AV canal GFP⁺ cells per embryo were low (typically 0.6 – 0.9 thousand per embryo), 5000 cells per sample were gated and used for RNA isolation. RNA was subjected to 2 rounds of linear amplification and biotin labeling. Within each group (Figure 5A) 6 labeled antisense RNA samples were hybridized to separate Illumina Mouse-Ref-6 BeadChips (Figure 4B). Present and absent calls were calculated, transcripts were counted as differentially expressed at a p-value, adjusted for multiple testing, of <0.01. Microarray data have been submitted to the Gene Expression Omnibus database [http://www.ncbi.nlm.nih.gov/projects/geo] under series GSE13614.

Genes known to be specific to the AV myocardium, including Tbx3 itself, were highly enriched in both E10.5 and E17.5 AV myocardial groups (Fig. 5C). Also the array data regarding Tbx20 were consistent with its initial broad and later AV canal-enriched pattern. Genes known to be specific to working myocardium were strongly enriched in working myocardium at both stages (Figure 5C). These data validate the microarray analysis,
and indicate it efficiently identified genes differentially expressed between the developing AV node and age-matched working myocardium. Mutations in *Nkx2-5*, *Tbx5* and *Gata4* result in AV conduction defects or structural AV canal defects in human and mouse (Bruneau, 2008). However, they were not enriched in the AV canal.

**Diverging phenotypes of the developing AV node and working myocardium**

Of 46643 transcripts present on the array, 14708 transcripts (32%) gave a signal above threshold and were considered to be expressed. Of these, 1259 (3%) were over- and 856 (2%) were underrepresented in the E10.5 AV canal compared to E10.5 WM (Figure 5A, B). In the E17.5 AV node and E17.5 WM, 15921 transcripts were expressed, similar to E10.5. However, the number of differentially expressed enrich genes strongly increased. 3182 (7%) transcripts were upregulated, and 3370 (7%) were under-represented in the E17.5 AV node relative to E17.5 WM, revealing an evident divergence of gene profiles of the differentiating AV node and complementary working myocardium. To assess this deviation in gene profiles we used the GLOBAL test (Goeman et al., 2004). Using standardized test-conditions, we identified 124 differentially expressed Gene Ontology (GO)-categories at E10.5, and 432 terms at E17.5. The test-conditions are shown in the online data supplement. The extensive lists of differentially expressed GO terms will not be discussed here, but are shown in the online data sets IIA and IIB. These data confirmed divergence of gene expression profiles and phenotypes between maturing AV nodal and working myocardium.

**Node-specific gene expression increases during development**

To evaluate AV nodal gene expression during development independently of developmental changes in working myocardial gene expression, the expression profile data of the E10.5 WM cell population was used as common reference for the three other data sets. We calculated genes differentially expressed between E10.5 WM and E17.5 AV nodal and E10.5 WM and E17.5 WM, respectively, and combined these data with the values found comparing both E10.5 groups. Whereas 2115 transcripts were expressed differentially between E10.5 WM and AV canal myocardium, the difference between E10.5 WM and E17.5 AV nodal myocardium had increased to 11388, implying an increase in differential expression of 9273 transcripts in the maturing AV nodal cells (Figure 6A). Of these, 4611 transcripts appeared differentially expressed between E10.5 WM and E17.5 WM as well, indicating these were not node-specific, but characteristic for myocardial maturation in general (Figure 6B). To gain insight into nodal differentiation, we assessed transcripts differentially expressed between E10.5 WM and E10.5 and/or E17.5 nodal tissue (Figure 6C), excluding those transcripts associated with myocardial maturation. In the E10.5 AV canal, 1092 transcripts were differentially expressed, in the E17.5 AV node 5856. Of the transcripts differentially expressed at E10.5, 249 (23%) transcripts were differentially expressed only early in
Figure 6. Evaluation of node-specific gene expression during development. (A) Venn diagrams showing transcripts (differentially) expressed in E10.5 AV canal (N10.5) and E17.5 AV node (N17.5) myocardium compared to common reference E10.5 WM (W10.5). Expression levels in E17.5 working myocardium (W17.5) were not taken into account. (B) Transcripts which, relative to W10.5, were differentially expressed in N10.5 and/or N17.5 (relative to W10.5), but not W17.5. Overlap reveals genes involved in node-specific expression throughout development (See Figure 5A, III). Boxes specify the number of transcripts over- or under-represented relative to W10.5. (D) Functional classification, using GO terms, of transcripts present on the microarray, and of differentially expressed transcripts of node-specific groups mentioned in Figure 6C. By converting the transcript-list into DAVID gene IDs which are unique per gene entry, potential redundancy was removed.
development, whereas 77% (843) were still differentially expressed at E17.5. Together, these data indicate that the AV node differentiates considerably during development, but that the late fetal AV node largely maintains the AV canal program acquired at E10.5.

To gain insight into the 3 groups of differentially expressed genes in the developing AV node, we used the Database for Annotation, Visualization and Integrated Discovery (http://david.abcc.ncifcrf.gov) bioinformatics tool to functionally categorize the transcripts, using GO annotations (Figure 6D). Found percentages were compared to the relative presence of specific GO terms on the Illumina Mouse-Ref-6 BeadChip (Figure 6D). Among transcripts enriched in both the AV canal and AV node, GO categories associated with cell differentiation, transcription and nervous system development were overrepresented. Among nervous system development, neurotrophic factors Nr1, Ntng1, Ntnt2l, Gdnf, Slit2 and Rtn4, the receptors for these neurotrophic factors Unc5b, Unc5c, Ret, and Robo2, and several semaphorins, including Sema3f, Sema3b, Sema4c, Sema4g and Sema6a, were found to be enriched in the AV canal and/or node. Contamination of nervous tissue at E10.5 can be excluded as the innervation of the heart has not yet taken place at this stage. Therefore, the latter finding should be attributed to shared molecular properties between developing nodal myocardium and nervous tissue. In addition, structure-related GO term cytoskeleton was up in both the AV canal and AV node. Among transcripts expressed at a lower level in nodal tissue at both stages, GO terms associated with contraction, energy and electrophysiology were overrepresented (muscle development, mitochondrion, cytoskeleton, contractile fiber, cell junction, ion channel activity). These processes and structures are typically more developed in the working myocardium (de Haan, 1961), further elaborating the notion that while AV canal cells substantially differentiate to form AV nodal cells, their primitive, non-working properties are maintained.

Transcripts differentially expressed exclusively in the E10.5 AV canal define its embryo-specific characteristics and might reveal important information regarding the initiation of conduction-system development. Among these, GO categories cell differentiation, muscle development and cytoskeleton were overrepresented both in the group of AV enriched and in the group of AV reduced transcripts (Fig. 6D). These representations highlight both the further differentiation of cardiac muscle during development, and the divergence in muscular phenotypes between both myocardial subpopulations.

The AV canal / node population could be contaminated with co-purified mesenchymal cells of the AV cushions / fibrous insulation that are in close association with these nodal cells. Resorting of purified GFP+ cells revealed approximately 85-90% pure cell populations. Indeed, in both E10.5 and E17.5 AV nodal groups, mesenchymal transcripts like Postn (Kruzynska-Frejtag et al., 2004) that are highly expressed specifically in the cushion mesenchyme were found to be significantly enriched, whereas the signals for these transcripts was low or absent in both WM groups. It is difficult to estimate contamination because of most transcripts the precise spatial distribution has not been established.
Chapter 4

Genome-wide profile of ion channel families expressed in AV nodal tissue

The AV myocardium has highly specific electrophysiological properties. To gain insight into the associated gene profiles, we examined the ion channel expression profiles of the E10.5 and E17.5 AV nodal myocardium relative to age-matched working myocardium (Table 1; Supplementary Table 3). In addition to previously established differences (Schram et al., 2002; Marionneau et al., 2005), we noted several novel channel transcripts to be enriched in the AV node or working myocardium. Of voltage-gated Na\(^{+}\) (Nav) channels and accessory genes, α-subunit Scn7A (Nav2.3) and β-subunit Scn2B (Navβ2) were higher expressed in the E10.5 AV canal and E17.5 node, whereas sodium channel modifier gene 1 (Scnml1) was enriched only in the E17.5 AV node. In working myocardium, the prominent cardiac α-subunit Scn5A (Nav1.5) was higher expressed at both developmental stages (Nerbonne and Kass, 2005), as well as fibroblast growth factor Fgf12, a growth factor recently shown to interact with Scn5A and modulate its properties (Liu et al., 2003). Additionally, Scn10A (Nav1.8), another neuronal type Nav α-subunit, was overrepresented in the working myocardium.

Of pore-forming voltage-gated calcium (Cav) channel α-subunits, Cacna1g (Cav3.1) was higher expressed in AV nodal cells at both stages, Cacna1c (Cav1.2), Cacna2d1 (Cava2δ1) and Cacna2d2 (Cava2δ2) were picked up at comparable levels in nodal and working myocardial tissue, and Cacna1h (Cav3.2) was expressed at comparable levels throughout E10.5 myocardium, but at E17.5 was enriched in atrial myocardium. Of accessory β-subunits, Cacnab1 (Cavβ1) was overrepresented in the E17.5 AV node, Cacnab2 (Cavβ2) was up in working myocardium at both stages. Other Cav related proteins were not detected above threshold. Of genes involved in Ca\(^{2+}\) homeostasis, Calmodulin3 (Calm3) was expressed at high levels throughout the myocardium, yet Calmodulin1 was not detected. Interestingly, of the Calcium/Calmodulin-dependent protein kinase family, several genes were enriched in the node, others in working myocardium. In addition to transient receptor potential (Trp) cation channels Trpc2, Trpc6, Trpm8, known to function in calcium homeostasis, several novel members of this family were enriched in AV nodal (Trpc1, Trpm5, Trpv4) or working myocardial tissue (Trpm2). Inositol 1,4,5-triphosphate receptor 3 (Itpr3), previously associated with pacemaker activity in differentiating cardiomyocytes, (Kapur and Banach, 2007) and Trpv4 were enriched only in the AV canal. Ankyrin-1,-2 and -3, ATPases, calsequestrin-2 (Casq2), histidine rich calcium binding protein (Hrc), phospholamban (Pln) and ryanodine receptor-2 (Ryr2) were all (much) higher expressed in working myocardium.

Among the voltage-gated potassium (Kv) channel family, α-subunits Kcna6 (Kv1.6), Kcnc4 (Kv3.4) and Kcnh1 (Meag) were enriched moderately in E10.5 AV canal and highly in E17.5 AV node. Auxiliary Kv channel subunits Kcne4 (MiRP), Kcne5 (Mink), Kcnip3 (KChIP3) and Pias3 (KChap) were enriched in the E17.5 AV node. The genes encoding α-subunits Kv1.4, Kv3.2, Kv4.2, Kv4.3, Kv6.2, erg6, Kcnq1 (KvLQT1) and channel modifier
genes *Dpp6* and *Kcmf1* (*Pmcf*) predominated in working myocardium, of which several already at E10.5. Of the inwardly rectifying K+ (Kir) channel family, α-subunits Kir3.1, Kir2.4, Kir6.1, Kir6.2 and Kir2.2 were higher expressed in the working myocardium. Of 3 differentially expressed members of a novel type of K+ pore-forming subunits (K2P), Kcnk2 (*TREK-1*) and Kcnk6 (*Twik2*) predominated in nodal (Kcnk2 only at E10.5) and Kcnk3 (*TASK-1*) in working myocardial tissue. Also, 3 calcium-activated channels were differentially expressed, of which Kcnma1 and Kcnn4 were enriched in the AV node. Other channel, accessory subunit and gap junctional transcripts are shown in Supplementary Table 3.
Discussion

Assessing the unique gene profile of the developing AV node myocardium

The adult AV node is a morphologically complex structure with heterogeneous molecular and functional characteristics containing not only cardiomyocytes, but also mesenchymal cells, fibroblasts, smooth muscles, innervating neurons and extracellular matrix. Here, we identified novel Tbx3 sequences that in the heart are exclusively active in the AV canal- and node myocardium, allowing their selective fluorescent labeling. The specificity of GFP expression was confirmed by detailed expression analysis and patch-clamp analyses of fluorescent cells. Using FACS we obtained purified AV myocardial cell populations largely free of other cell types, and their expression profiles were compared to those of age-matched working myocardial cells obtained from NppaBAC336-Egfp mice (Horsthuis et al., 2008). Some contamination of other cells-types, such as AV mesenchymal cells, was observed in the purified AV myocardial populations, but these appeared minor. All transcripts that we are aware of to be enriched in either AV- or (developing) working myocardium were found to be enriched in the corresponding cell populations (Figure 5 and Table 1). Furthermore, genes associated with mitochondria and contractile fibers were underrepresented in both the AV canal and AV node, whereas Nervous system development (especially a subclass of neurotrophic genes and semaphorins) was prominently overrepresented in both the AV canal and AV node. These data not only confirm previous findings (de Haan, 1961; Virágh and Challice, 1977a; Virágh and Challice, 1977b; Moorman et al., 1998; Walls, 1947), but provide a much more elaborate picture of the transcriptional landscape of the developing AV node and working myocardium. Taken together, we have generated a potentially valuable transgenic tool to study AV node development and function in vivo.

To gain insight into the molecular underpinning of the physiologically complex AV node, we analyzed the expression profile data sets for expression of 'electrophysiological' transcripts (ion channel, accessory subunit, etc). Previous expression studies of the adult AV node have provided valuable data on the expression distribution of those genes (Marionneau et al., 2005). However, those were performed on dissected adult tissues contaminated with adjacent working myocardium and the non-myocardial cell populations, potentially affecting the observed expression profiles. Furthermore, they examined expression of 82 pre-defined genes (Marionneau et al., 2005). The microarray approach presented here largely confirmed these findings, and in addition identified a considerable number of novel cardiac transcripts enriched in either AV nodal or working myocardium that may be important for understanding AV node function (Table 1 and Supplementary Table 3).

The origin and modular development of the AV node

Histological studies and marker analyses revealed that the conduction system (sinus node, AV node, AV bundle and bundle branches) is formed from a contiguous network of myocardial cells distinctive from the surrounding atrial and ventricular (working)
myocardium (Virágh and Challice, 1977a;Virágh and Challice, 1977b;Moorman and Christoffels, 2003;Gourdie et al., 2003). This network can be discriminated already early in development by poorly developed sarcomeres and sarcoplasmatic reticulum, high automaticity, low proliferation rate, sparse mitochondria, selective expression of markers (e.g. Glu2, HNK-1, Tbx3, minK-lacZ, CCS-lacZ) and absence of high-conductance gap junctions (e.g. Cx43) (de Haan, 1961;Virágh and Challice, 1977a;Virágh and Challice, 1977b;de Jong et al., 1992;Moorman et al., 1998). These properties, which are controlled by Tbx3 (Hoogaars et al., 2007;Bakker et al., 2008) and other factors, are largely maintained in the adult components. The existence of this contiguous precursor network with shared properties suggests a common regulatory mechanism for conduction system gene expression and formation. On the other hand, arguments in favor of a modular composition have been put forward as well. During mid-fetal stages, the initially 'nodal' AV bundle acquires rapid conduction and Cx40 expression (Chuck et al., 1997;Bakker et al., 2008). In Nkx2-5/Tbx5 and Tbx5/Id2 double heterozygous mutants and Tbx3 mutants, which develop severe AV bundle defects, the AV node may be less affected (Moskowitz et al., 2007;Bakker et al., 2008), suggestive for differential sensitivity of these components to loss of transcription factor function. In the present study, we found that 160 kbp of Tbx3 and flanking sequences drive expression highly selectively in the AV canal of embryos, and AV node of adults, but not in the Tbx3+ AV bundle or sinus node. These data unambiguously demonstrate that separable regulatory sequences and regulatory pathways (regulatory modules) control gene expression and formation of the AV node, AV bundle and other conduction system components.

The AV node may be derived from the embryonic AV canal or AV ring, or may form de novo from other sources (Virágh and Challice, 1977a;Virágh and Challice, 1977b;Arguéllo et al., 1988). From E9 onwards, the expression of GFP was observed selectively in the myocardium of the AV canal, and gradually became confined to the AV node and anterior node-like structure (Blom et al., 1999). Further, the expression profile of the embryonic AV canal showed a large overlap with the profile of the late fetal AV node, whereas both profiles were found to be consistently distinctive from both E10.5 and E17.5 working myocardium. These findings indicate early segregation of AV canal cells and adjacent developing working myocardial cells from E10.5 or earlier, arguing the AV node arises from myocardium in the dorsal wall of the AV canal as proposed by Viragh and Challice (Virágh and Challice, 1977a;Virágh and Challice, 1977b).

**Specialization of the AV nodal myocardium during development**

The embryonic AV canal and the derived AV node maintain many properties (see above) found already in the embryonic heart tube, whereas the surrounding myocardium acquires working myocardial properties (Moorman and Christoffels, 2003). These observations have led to the idea that suppression of (working myocardial) differentiation in the AV canal domain is an important contributing mechanism to AV node formation, an idea that gained
strong support by the functional identification of suppressors Tbx2 and Tbx3 in the AV canal and conduction system (Moorman and Christoffels, 2003). However, the AV myocardium itself is likely to also specialize during development (Milan et al., 2006), an aspect that has not been thoroughly examined thus far. When comparing the AV gene expression profiles at an early (E10.5) and late (E17.5) time point of development with the common reference, 249 transcripts were found to be differentially expressed only at E10.5. These are interesting candidates potentially involved in the initiation and early differentiation of the AV node. 77% of transcripts (843 transcripts) differentially expressed at E10.5 still are differentially expressed in the fetal AV node, indicative for maintenance of properties during AV node development. On the other hand, around 5 thousand transcripts not differentially expressed at E10.5 were differentially expressed in the E17.5 AV node, indicating that the AV node undergoes substantial specialization during development.

**Acknowledgments**

We thank Berend Hooibrink for excellent technical support, Corrie de Gier-de Vries and Wim TJ Aanhaanen for their contributions, Daniel J Garry and Amanda M Masino for providing protocols, and Marian van Roon of the AMC GGM facility for generating transgenic mice. This work was supported by grants from the Netherlands Heart Foundation (96.002 to V.M.C. and A.F.M.M.); European Community’s Sixth Framework Programme contract ('HeartRepair') LSHM-CT-2005-018630 to AMC (V.M.C., A.F.M.M.) and LUMC (P.A.C.H.).
References


Supplementary data

Expanded materials and methods

Transgenic mice

-6/+2mLacZ construct
To generate the -6/+2mLacZ construct, an nlsLacZ-pA cassette was placed at the translation start site of Tbx3 in an 8 kbp fragment from the endogenous Scal restriction site at 6 kbp upstream to the endogenous PshA1 site at 2 kbp downstream of the translation start site. Vector sequences were removed before introduction into the mouse genome.

Tbx3BAC-Egfp construct
The two-step BAC modification protocol previously described by Shiaoching Gong and Nathaniel Heinz (Gong et al., 2002), consists of two homologous recombination steps. After both the co-integration and the resolution step correct recombination was verified by Southern blot using a hybridization probe against Egfp. The BAC-Egfp constructs were purified using a CsCl gradient following a protocol also kindly provided by Shiaoching Gong and Nathaniel Heinz. Constructs were linearized before introduction into the mouse genome. To generate transgenic mouse lines, constructs were injected into pronuclei of zygotes of FVB mice and these were re-implanted into pseudo-pregnant foster mothers by use of standard techniques. Animal care was in accordance with national and international guidelines.

Non-radioactive in situ hybridisation, β-galactosidase activity detection and immunohistochemistry
Embryos were fixed in 4% formaldehyde, embedded in paraplast and sectioned at 10 µm for immunohistochemistry and in situ hybridization on sections. Whole mount in situ hybridization, in situ hybridization on sections and whole mount β-galactosidase activity staining were performed as described previously (Moorman et al., 2001;Franco et al., 2001). For immunohistochemistry GFP rabbit polyclonal primary antibody (1:250; Santa Cruz) was used. The secondary antibody used was Alexa 568 goat-anti-rabbit (1:250; Invitrogen).

Electrophysiology
Cell isolation. Hearts isolated from E17.5 heterozygous BACTbx3-Egfp+ embryos were immersed in a standard solution containing (mmol/l) 140 NaCl, 5.4 KCl, 1.8 CaCl2, 1.0 MgCl2, 5.5 glucose, 5 HEPES; pH7.4 (NaOH). After removal of the atria, fluorescent cells of the atrioventricular (AV) region were visualized with a stereomicroscope equipped for GFP detection. The fluorescent AV regions of 10 embryo’s were dissected and pooled. Cells were
dissociated in an enzyme mix (Gibco 14170-070 with pancreatin, 60 mg/ml and trypsin 1.25 mg/ml) in a multiple rotate-harvest procedure, using a rotator and water bath at 37°C. To get rid of cardiac fibroblasts cells were incubated for 2 hours, next myocytes were plated on collagen coated cover slips and cultured for 24 hours.

**Data acquisition.** Action potentials (APs) and hyperpolarizing-activated current (I<sub>f</sub>) were recorded at 36±0.2°C from single BACTbx3-Egfp<sup>+</sup> and BACTbx3-Egfp<sup>-</sup> cells with the amphotericin-B perforated patch-clamp technique using an Axopatch 200B patch-clamp amplifier (Molecular Devices Corporation, Sunnyvale, CA, USA). Superfusion solution contained (mmol/l): 140 NaCl, 5.4 KCl, 1.8 CaCl<sub>2</sub>, 1 MgCl<sub>2</sub>, 5.5 glucose, 5 HEPES; pH 7.4 (NaOH). Pipettes (3-4 MΩ, borosilicate glass) were filled with solution containing (mmol/l): 125 K-gluc, 20 KCl, 5 NaCl, 0.22 amphotericin-B, 10 HEPES; pH 7.2 (KOH). APs were low-pass filtered with a 10-kHz cut-off frequency, and digitized at 25-kHz; I<sub>f</sub> recordings were both filtered and digitized at 5-KHz. Membrane capacitance (C<sub>m</sub>) was estimated by dividing the decay time constant of the capacitive transient in response to 5-mV hyperpolarizing voltage-clamp steps from -40 mV by the series resistance (R<sub>s</sub>). Potentials were corrected for liquid junction potential, and C<sub>m</sub> and R<sub>s</sub> were compensated for >80%. For voltage control, data acquisition, and analysis, custom-made software was used.

**Stimulation protocols.** (Supplementary Table 2) In cells without pacemaker activity, APs were elicited at 2-Hz by 3-ms, 1.5× threshold current pulses through the patch pipette. We analyzed cycle length, maximal diastolic potential (MDP), diastolic depolarization rate (DDR, measured over the 100-ms time interval starting at MDP + 1 mV), maximal upstroke velocity (dV/dt<sub>max</sub>), maximal AP amplitude (APA), and AP duration at 20, 50, and 90% repolarization (APD<sub>20</sub>, APD<sub>50</sub> and APD<sub>90</sub>). Parameters from 10 consecutive APs were averaged. I<sub>f</sub> was measured by 500-ms hyperpolarizing voltage clamp steps from a holding potential of ~40 mV. I<sub>f</sub> was normalized to C<sub>m</sub>.

**Statistics.** Results are expressed as mean±SEM. Data are considered different if p<0.05. The Mann-Whitney U test, unpaired t-test, or Fisher’s exact test was used if appropriate.

**Fluorescent-activated cell sorting and RNA handling**

Hearts from stage-matched embryos were removed, pooled, and single-cell FACS samples were prepared using a protocol described by Masino and Garry (Masino et al., 2004). Cells were sorted on a FacsAria flow cytometer (BD Biosciences) using FacsDIVA software. Samples from wild-type age-matched hearts were used for gating. Samples were gated to exclude debris and cell clumps. Per sample 5000 live-gated cells were collected directly in RAA Nucleospin RNA cell lysis buffer and RNA was isolated following the protocol of the manufacturer (Machery-Nagel). RNA integrity was checked on an Agilent Bioanalyzer 2100, using RNA 6000 pico chips. Samples were subjected to 2 rounds of linear amplification, using the MessageAmp II aRNA Amplification Kit (Ambion) for the first round, and the
Illumina TotalPrep RNA Amplification Kit (Ambion) for the second round and biotin-16-UTP labelling.

**Microarray analysis**

Labeled RNA was hybridized to Illumina MouseRef-6 BeadChip following the manufacturer’s instructions (Illumina Inc.). Beadstudio software was used to assess the individual array quality. Intensity values were averaged per bead type and subsequently normalized using VSN in R (Huber et al., 2002). Present/absent call calculation was based on more than 300 negative controls per array. Analysis of differentially expressed genes was done with LIMMA package in R (Smyth et al., 2004). P-values were adjusted for multiple testing using the method proposed by Benjamin and Hochberg. Transcripts were counted as differentially expressed if the adjusted p-value was <0.01.

**Bioinformatics**

*Settings GLOBAL test determining global (differential) expression patterns of functional groups of genes*

A Gene Ontology (GO) functional category is considered to be significantly associated with the difference in genotypes tested when all of the following selection criteria are met.

Per GO category:

1) at least 10, maximum 75 genes
2) at least 5 genes with an influence >5
3) at least 25% of genes for N>W with influence >5, or at least 25% of genes for W>N with influence >5, or both
4) BY-FDR <= 0.1

*Geneplots of differentially expressed GO categories:*

The bar height for each of the genes in the geneplots indicates the standardized influence of each gene represented in the specific GO term. The scales in the individual bars represent the number of standard deviations that the gene influence exceeds the null hypothesis that the gene is not associated with the difference in phenotype between N and W for the specific time point. Genes with red bars have higher expression levels in W, genes with green bars have higher expression levels in N. The blue dashed line indicates the critical influence value of 5 that was used in the selection of the GO terms. For practical reasons, geneplots have not been included here. For geneplots of differentially expressed GO categories please contact the corresponding author (e-mail: v.m.christoffels@amc.uva.nl).
**Supplementary Table 1.** Overview of expression patterns of the endogenous Tbx3 gene and the 160 kbp BACTbx3-Egfp construct, E9.5-E17.5. +, expression; -, no expression; +/-, incomplete expression.

<table>
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<th><strong>Egfp</strong></th>
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<tr>
<td>sinus node</td>
<td>+</td>
</tr>
<tr>
<td>AV canal</td>
<td>+ +/-</td>
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<tr>
<td>AV cushions</td>
<td>+</td>
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<tr>
<td>AV node</td>
<td>+</td>
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<tr>
<td>AV (His) bundle</td>
<td>+ -</td>
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<tr>
<td>second heartfield</td>
<td>+/- -</td>
</tr>
<tr>
<td>OFT cushions/valves</td>
<td>+ -</td>
</tr>
<tr>
<td>cardiac neural crest</td>
<td>+ +</td>
</tr>
<tr>
<td>cardiac plexus</td>
<td>+ +/-</td>
</tr>
<tr>
<td>mandibula mesenchyme</td>
<td>+ +</td>
</tr>
<tr>
<td>maxilla mesenchyme</td>
<td>+ +</td>
</tr>
<tr>
<td>whisker</td>
<td>+ +</td>
</tr>
<tr>
<td>teeth</td>
<td>+ +</td>
</tr>
<tr>
<td>tongue</td>
<td>+ +</td>
</tr>
<tr>
<td>optic vesicle (dorsal region)</td>
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<tr>
<td>otic vesicle</td>
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<td>infundibulum</td>
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<td>thyroid</td>
<td>+ +</td>
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<tr>
<td>hypoglossal cord</td>
<td>+ +</td>
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<tr>
<td>laryngeal groove mesenchyme</td>
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<tr>
<td>laryngeal groove epithelium</td>
<td>- +</td>
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<tr>
<td>trachea mesenchyme</td>
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<td>diaphragm</td>
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<td>liver</td>
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Supplementary Table 2. Action potential properties of single BACTbx3-Egfp-positive and BACTbx3-Egfp-negative cells

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<tr>
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<th>GFP(^-) cells</th>
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<td>n=4</td>
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<td>n=5</td>
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<tr>
<td>cycle length (ms)</td>
<td>472±138</td>
<td>–</td>
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<tr>
<td>MDP (mV)</td>
<td>-58.9±5.1</td>
<td>-74.2±2.7(^*)</td>
</tr>
<tr>
<td>DDR(_{100}) (mV/s)</td>
<td>55±27</td>
<td>–</td>
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<tr>
<td>dV/dt(_{\text{max}}) (V/s)</td>
<td>27±14</td>
<td>111±30(^*)</td>
</tr>
<tr>
<td>APA (mV)</td>
<td>97±7</td>
<td>123±5(^*)</td>
</tr>
<tr>
<td>overshoot (mV)</td>
<td>38±10</td>
<td>48±5</td>
</tr>
<tr>
<td>APD(_{20}) (ms)</td>
<td>27.8±7.1</td>
<td>4.4±1.0(^*)</td>
</tr>
<tr>
<td>APD(_{50}) (ms)</td>
<td>61.2±8.4</td>
<td>14.8±5.1(^*)</td>
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<tr>
<td>APD(_{90}) (ms)</td>
<td>98.6±13.0</td>
<td>44.8±7.4(^*)</td>
</tr>
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Data are mean±SEM; n, number of cells; MDP, maximal diastolic potential; DDR\(_{100}\), diastolic depolarization rate over the 100-ms time interval starting at MDP; dV/dt\(_{\text{max}}\), maximal upstroke velocity; APA, action potential amplitude; APD\(_{20}\), APD\(_{50}\), and APD\(_{90}\), action potential duration at 20, 50, and 90% repolarization. \(^*\)p<0.05.
**Supplementary Table 3.** Expression profiles of all electricity-associated transcripts on the microarray, categorized in functional subclasses.

<table>
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<th>E17.5 Fold higher</th>
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**Sodium channel and auxiliary subunits, modifier genes**

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E10.5 and E17.5 AV canal/node (N) compared with age-matched working myocardium (W). N>W, higher expression in N than in W (p<0.01); W>N, higher expression in W than in N (p<0.01); N=W, not differentially expressed between N and W (p>0.01), expression level above threshold in N and/or W; -, not differentially expressed between N and W (p>0.01), expression level not above threshold in N and W; *, Expressed above threshold, some probes of gene higher signal in N, some in W

References supplementary data


Summary

Within 3 weeks of conception in humans (comparable with mouse embryonic day E8.5), it is already possible to identify the beating heart. At this early stage, the heart is a tube-like structure, which contracts in a peristaltoid fashion, pumping the blood from the venous to the arterial pole. Its wall consists of an inner endocardial layer and an outer myocardial layer, the latter containing muscle cells sharing important ‘embryonic’ characteristics, among which high automaticity, low contractility and low conduction velocity. Subsequent to its formation the tube starts to loop to the right, and at specific places on the outer curvature of the looping heart a chamber-specific gene program is turned on, marking the initiation of the development of the atrial and ventricular chambers. The myocardium of the newly evolving atria and ventricles share typical ‘working myocardial’ characteristics, i.e. high contractility, high conduction velocity and low automaticity, making these chambers ideally equipped to pump the blood ahead in the now rapidly growing embryo. The rest of the looping heart tube, that is the inflow tract, atrioventricular canal (the embryonic structure connecting the developing atria and ventricles), inner curvature and outflow tract, does not differentiate into chamber myocardium, but maintains its embryonic phenotype for a longer period. From these structures the nodal parts of the conduction system, still characterized by the previously mentioned embryonic characteristics, will develop. The inflow tract gives rise to the sinus node, whereas the atrioventricular node develops from the back wall of the atrioventricular canal. During subsequent steps of embryonic development, by an interplay of additional steps of differentiation, cell growth, proliferation and recruitment of cells to the cardiac lineage, the 4-chambered heart is formed, containing numerous phenotypically different cells. In the mature heart, the atria and ventricles are fully septated, the pulmonal and the systemic circulation functioning in parallel. The heart contains valves ensuring efficient unidirectional blood flow, and the highly specialized cardiac conduction system ensures sequential, synchronized contractions of the atria and ventricles. On a molecular level many decisions have to be made before embryonic cardiac cells develop in a correct pattern from their primitive to their mature terminally differentiated state. This thesis focuses on the deciphering of some of the molecular regulatory networks, which orchestrate the decision of an embryonic myocardial cell to become a myocardial cell of the atrioventricular node or chamber.

Many of the decisions taken in (cardiac) morphogenesis are enforced by a complex interplay of transcription factors and signaling molecules. Transcription factors are proteins that are able to bind to specific sequences of DNA. When they bind, together with other transcription factors, signaling molecules and co-factors, they turn on, or turn off, the transcription of downstream target genes. These target genes, in turn, induce the changes in cellular differentiation. Whereas transcription factors are expressed intracellularly and direct their action upon the cell in which they are expressed, signaling molecules are excreted
extracellularly. These proteins act upstream of a ‘signal transduction pathway’. Upon excretion, signaling molecules bind to the extracellular part of highly specific trans-membrane receptors present only on target cells. Following binding, the signal is transduced intracellularly, as the intracellular part of the activated receptor turns on an intracellular signaling cascade. Eventually, this intracellular signaling cascade results in proteins shuttling from the cytoplasm into the nucleus of the target cell, where – just like transcription factors – they often bind to specific regulatory sequences of DNA, thereby activating or repressing the transcription of downstream target genes. In heart development, the T-box family of transcription factors plays a pivotal role as will be discussed in this thesis. Several signal transduction pathways play a role in the morphogenesis of the heart. Of these, several members of the subfamily of bone morphogenetic proteins (Bmp)s, which belongs to the transforming growth factor β superfamily, are known to closely interact with T-box transcription factors. The Bmp subfamily consists of more than 20 members, of which, in this thesis, Bmp2 – and the intracellular molecules Smad1, 5, 8 and 4, which shuttle into the nucleus upon Bmp activation - will be discussed in detail. *Nppa*, the gene encoding atrial natriuretic factor (ANF), of which our study of its regulatory DNA sequences is described in Chapter 2, is a typical example of a target gene.

**Chapter 1** reviews recent insights into some key morphogenetic processes underlying early cardiac development and the implications of this new knowledge for our thinking about the genesis of congenital heart defects. First, we discuss current understanding of patterning along the cranio-caudal, dorso-ventral, and left-right axes of the initially straight heart tube, explaining, among other malformations, why in the heart only the atria can be truly isomeric. Then, recent insights into the formation of the atrial and ventricular chambers, and their underlying transcriptional networks, are discussed. In many cardiac textbooks still the incorrect segmental model of cardiac formation is described, in which it was inaccurately assumed that already in the early heart tube all progenitors of the building blocks of the mature heart are present as a linear array of transverse precursor segments. Here, we review the up-to-date ballooning model of chamber formation, the first model which clearly explains how an initially solitary tube, with laminar flow through a single lumen, can transform into a 4-chambered heart in which the pulmonary and systemic circulations work in parallel. Finally, we discuss novel information on the origin of the cells that are added to the heart after formation of the initial tube, and their fate during further development.

In **Chapter 2** we explore the regulatory DNA sequences driving expression of *Nppa*, the gene encoding atrial natriuretic factor (ANF), during development and disease. Before birth *Nppa* is expressed specifically in the evolving atrial and ventricular chambers, marking their development. After birth the gene is silenced in the ventricles, where, in case of cardiac hypertrophy or heart failure, it is reactivated again as part of a larger ‘fetal’ gene program. To get insight into the regulatory sequences of *Nppa*, we generated several transgenic mice, all of them carrying a different transgenic construct integrated into their genome, in which a unique sequence of regulatory DNA, cloned from the endogenous *Nppa* locus, drives a reporter gene.
(green fluorescent protein (GFP), LacZ or Cre). Then we induced heart failure in the reporter mice by crossbreeding them with another transgenic mouse model, in which the yeast transcription factor Gal4 is expressed throughout the myocardium, causing aspecific cardiomyopathy in males three weeks after birth. In some of the reporter mice, cardiac hypertrophy was induced by aortic banding, a procedure in which the ascending aorta is ligated, causing a pressure-overload hypertrophy model. Comparison of the expression patterns of the reporter genes in the diverse transgenic mice, both in embryos and in diseased hearts, with the expression pattern of endogenous Nppa, then provided insight into the potency of the diverse regulatory DNA sequences. From previous studies in transgenic mice, in which only short regulatory sequences were tested, it had been concluded that a proximal sequence of 0.7 kbp (kilo basepairs) contains all regulatory modules necessary to drive correct expression before birth. In those studies, the sequences driving Nppa expression during disease had never been located. By combining data from reporter mice carrying short regulatory fragments randomly integrated into the genome, with data from mice harboring a short fragment that was knocked in as a single copy into the genome, and data from reporter mice carrying very large regulatory fragments (up to 200 kbp in bacterial artificial chromosomes (BACs)), we came to very different conclusions. We show that, to drive correct ventricular activity during development, the proximal Nppa regulatory fragment lacks regulatory sequences. We then demonstrate that for correct ventricular expression before birth, at least two separate distally located activating sequences (enhancers) are required, one providing the correct pattern and one providing the correct fetal activity. Regulatory sequences driving Nppa expression during disease (stress response) might overlap or cooperate with those regulating the fetal expression pattern, but appear to function independently of the enhancer regulating ventricular activity before birth. Therefore, we conclude that ventricular Nppa activity before birth and reactivation during disease are regulated by separate regulatory sequences, and as a consequence by divergent transcriptional pathways. These findings implicate that all previous data on gene regulation obtained from transgenic mouse models using the short Nppa regulatory sequences, either during chamber development or cardiac hypertrophy and heart failure, should be re-evaluated. Additionally, they implicate that data on gene regulation obtained from fetal mouse models cannot automatically be applied on hypertrophy models, and vice versa.

In Chapter 3 we aim to deepen our knowledge on the molecular pathways underlying the localized formation of the chambers versus the atrioventricular canal. From previous studies, the T-box transcription factors Tbx2, Tbx3 and Tbx20, together with Bmp/Smad-signaling, already had been established to play pivotal roles in the development of the chambers (Tbx20) and the atrioventricular canal (Tbx2, Tbx3 and Bmp/Smad). However, their genetic hierarchies and spatial and temporal interplay had remained unclear. By investigating mice homozygous mutant for Tbx2, for Tbx3, or double mutant for both genes, we found that Tbx2 and its close relative Tbx3 are redundantly required and individually sufficient to specify atrioventricular myocardium, induce formation of the atrioventricular cushions and suppress chamber differentiation in the atrioventricular canal. Furthermore, by
studying reporter mice harboring diverse Tbx2-reporter constructs and transfection experiments in which the constructs were temporally introduced into cell cultures in vitro, we identified a Bmp/Smad-dependent enhancer conferring atrioventricular canal-restricted expression and Tbx20-dependent chamber suppression of Tbx2 in vivo. Then we show that the T-box transcription factor Tbx20 plays a dual role in the compartmentalization of the heart. Contrary to what had been hypothesized previously, Tbx20 appears to stimulate chamber differentiation independently of Tbx2, and at the same time suppresses the activation of Tbx2 in the developing chambers, thereby localizing Tbx2 activity to the atrioventricular canal. Unexpectedly, Tbx20 does not repress Tbx2 expression directly by binding to its regulatory DNA sequences, but indirectly by binding to activated Smad1 and Smad5, sequestering these proteins from Smad4. Since Tbx2 is activated in the atrioventricular canal by Bmp2 in a Smad-dependent manner and Smad4 is a necessary co-factor for nuclear translocation and transcriptional activation by Smad1/Smad5, by this indirect mechanism Tbx20 effectively inhibits the Bmp/Smad-mediated activation of the target promoter of Tbx2. In conclusion, our findings suggest that opposing regulation of Bmp signaling by Tbx20 and Tbx2 may underlie specification of the chambers and the atrioventricular canal, respectively.

In Chapter 4 we describe the generation of a novel reporter mouse, in which green fluorescent protein (GFP) is driven by 160 kbp of regulatory DNA sequences of the T-box transcription factor Tbx3. Whereas endogenous Tbx3 is expressed in the different components of the cardiac conduction system, i.e. the sinus node, the atrioventricular node, the atrioventricular bundle and the bundle branches, the reporter gene was expressed specifically in the developing atrioventricular node but not in the other components of the cardiac conduction system. This finding demonstrates that the components of the conduction system are driven by distinct regulatory sequences, and therefore separate transcriptional pathways. The reporter mice were then used to purify atrioventricular nodal cells at two development stages (embryonic E10.5 and fetal E17.5). Subsequently, the gene expression profiles of these purified nodal cells were compared with age-matched working myocardial cells (purified using transgenic mice described in Chapter 2) using microarray analysis, a technique allowing examination of the activity of more than 20,000 genes at the same time. We constructed a comprehensive list of genes associated with electrical impulse propagation, which were picked up specifically in the nodal or working myocardium. Many of these were ion channels, not previously associated with nodal or working myocardium and several not even with cardiac expression. Furthermore, the data revealed that the atrioventricular node and the working myocardium phenotypes diverge during development, but that the functional gene classes characterizing both subtypes are maintained. One of the repertoires identified in the atrioventricular node-specific gene profiles consists of multiple factors previously only associated with neural development, revealing shared characteristics between nodal and nervous system development. These data present the first genome-wide transcription profiles of the atrioventricular node during development, providing valuable information concerning its molecular identity.
Drie weken na de bevruchting is het al mogelijk een kloppend hart waar te nemen in het zich ontwikkelende menselijke embryo. Zo vroeg in de ontwikkeling is het hart een buisvormige structuur, die het bloed van de veneuze instroom naar de arteriële uitstroom pompt door middel van peristaltische contracties. De wand van de hartbuis bestaat uit een binnenste endocardiale laag en een buitenste myocardiale spierlaag. De cellen van de spierlaag worden cardiomyocyten genoemd en hebben een aantal belangrijke ‘embryonale’ eigenschappen gemeenschappelijk, waaronder hoge automaticiteit (de eigenschap van een cel om zelf een elektrische prikkel te kunnen genereren), lage contractiliteit (de mogelijkheid tot samentrekken), en een lage impuls-voortgeleidingssnelheid. Nadat de rechte hartbuis is gevormd begint deze uit te bochten naar voren en naar rechts. Rond dezelfde tijd wordt op specifieke plaatsen van de nu ontstane buitenste bocht van de hartbuis een specifiek genprogramma aangeschakeld, waardoor de eigenschappen van de cellen ter plaatse veranderen (differentiëren). Dit proces markeert het begin van de ontwikkeling van de boezems (atria) en de kamers (ventrikels). De spiercellen van de zich ontwikkelende atria en ventrikels hebben andere, typische ‘werk’ eigenschappen gemeenschappelijk, zoals hoge contractiliteit en hoge voortgeleidingssnelheid, maar lage automaticiteit. Met deze eigenschappen zijn de kamers perfect toegerust om het bloed met kracht voort te pompen in het nu snel groeiende embryo. De rest van het uitbochtende hart, namelijk het instroom gebied (inflow tract), het atrioventriculaire kanaal (de embryonale structuur die de atria met de ventrikels verbindt), de binnenste curvatuur (dat gedeelte van de hartbuis dat zich aan de binnenbocht bevindt) en het uitstroom gebied (outflow tract), differentieert niet tot kamer-spiercel, maar behoudt zijn embryonale eigenschappen gedurende een langere periode (zie ook figuur 3 in hoofdstuk 2, waarin bovenstaand proces aan de hand van een tekening wordt uitgelegd). Uit deze laatstgenoemde structuren ontwikkelen zich onder andere de knopen van het geleidingssysteem van het hart, welke ook in het volwassen stadium gekarakteriseerd worden door de eerder genoemde ‘embryonale’ spiercel-eigenschappen. De sinusknoop, die ontstaat in het instroom gebied, is een gespecialiseerde spiercel-structuur bij de instroom van de bovenste lichaamsader in het rechter atrium. Deze structuur heeft de hoogste automaticiteit van alle hartspiercellen en genereert de elektrische prikkel die het hart aanzet samen te trekken. Vanuit de sinusknoop wordt de elektrische impuls voortgeleid door de atriale wand, welke als reactie samentrekt, waarna de impuls terechtkomt in de atrioventriculaire knoop. De atrioventriculaire knoop ontstaat uit het atrioventriculaire kanaal. In het volwassen stadium is de atrioventriculaire knoop de enige myocardiale structuur die de atria, via de atrioventriculaire bundel, met de ventrikels verbindt. Verder zijn de atria en ventrikels elektrisch gescheiden door bindweefsel. De atrioventriculaire knoop heeft twee belangrijke eigenschappen. Ten eerste vertraagt de atrioventriculaire knoop de impuls voordat hij deze
doorgeeft aan de ventrikels, waardoor deze net iets later samentrekken dan de atria. Hierdoor profiteren de ventrikels optimaal van de extra instroom van bloed door de atria-contractie, en pompen dus een zo groot mogelijk volume per slag. Ten tweede heeft ook de atroventriculaire knoop een hoge automaticiteit, zodat, wanneer de sinusknoop uitvalt, de atroventriculaire knoop een prikkel kan genereren die de kamers aanzet samen te trekken. Gedurende hieropvolgende stappen van ontwikkeling wordt, door een samenspel van differentiatie, celgroei, celdeling, en toevoeging van cellen aan het hart, het vierkamerige volwassen hart gevormd. Dit bevat veel fenotypisch verschillende cellen. In het volwassen hart zijn het linker en rechter atrium, evenals de linker en rechter ventrikel, volledig van elkaar gescheiden door tussenschotten, waardoor de long- en de lichaams-bloedsomloop gescheiden functioneren. Het hart bevat kleppen die een efficiënte bloedstroom in één richting verzekeren, en het gespecialiseerde geleidingssysteem van het hart verzekert opeenvolgende, synchrone contractie van de atria en de ventrikels. Op moleculair niveau moeten veel beslissingen worden genomen voordat de embryonale hartcellen zich in een correct patroon ontwikkelen van hun primitieve tot hun terminaal gedifferentieerde staat. Dit proefschrift richt zich op het ontrafelen van sommige van de moleculaire regel-netwerken, die ten grondslag liggen aan de beslissing van een embryonale spiercel om zich te ontwikkelen tot een spiercel van de atroventriculaire knoop, dan wel tot een spiercel van de (atriale of ventriculaire) kamers.

Binnen één organisme bevat elke cel hetzelfde erfelijk materiaal (genoom of DNA). De reden dat elke cel toch andere eigenschappen heeft (een ander fenotype heeft), is dat in elke cel verschillende genen aan en uit staan. Als een gen aan staat, wordt deze afgelezen in RNA, een proces dat transcriptie wordt genoemd. Vervolgens wordt dit RNA omgezet in eiwit. Dit proces heet translatie. Alle eiwitten die aanwezig zijn in een specifieke cel bepalen samen het fenotype van deze cel, met andere woorden bepalen welke eigenschappen deze cel heeft. Binnen elke cel wordt zowel de transcriptie als de translatie nauwkeurig gereguleerd. Veel van de beslissingen die worden genomen gedurende de ontwikkeling worden op transcriptieniveau opgelegd door een complex samenspel van transcriptiefactoren en signaalmoleculen. Transcriptiefactoren zijn eiwitten die kunnen binden aan een specifieke DNA regel-sequentie. Elk gen heeft zijn eigen regulatoire sequenties, welke bestaan uit een specifieke volgorde van de bouwstenen van het DNA (basenparen). Als transcriptiefactoren binden, in samenspel met andere transcriptiefactoren, signaalmoleculen en co-factoren, schakelen ze de transcriptie van doelwit-genen aan of uit. De hieruit resulterende verhoogde of verlaagde aanwezigheid van doelwit-eiwitten in de cel induceert uiteindelijk de veranderingen in cellulaire differentiatie. Terwijl transcriptiefactoren intracellulair (binnen de cel) tot expressie komen en dus de differentiatie van de cel beïnvloeden waarin ze actief zijn, worden signaalmoleculen extracellulair (buiten de cel) uitgescheiden. Deze signaaleiwitten fungeren als meest ‘stroomopwaarts’ gelegen spelers van een ‘signaal-transductie pad’. Na uitscheiding binden de signaaleiwitten aan het extracelluliere gedeelte van specifieke transmembraanreceptoren die alleen aanwezig zijn op doelwit-cellen. Na binding wordt het signaal naar het binnenste van de cel (het cytoplasma) doorgegeven, doordat het intracellulaire
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gedeelte van de geactiveerde receptor een intracellulaire signaalcascade activeert. Uiteindelijk leidt deze cascade ertoe dat specifieke eiwitten vanuit het cytoplasma de celkern in migreren, waar ze – net als transcriptiefactoren – meestal aan specifieke DNA regel-sequenties binden, en zo de transcriptie van doelwit-genen activeren of onderdrukken.

Zoals wordt besproken in dit proefschrift, speelt de T-box familie van transcriptiefactoren een centrale rol tijdens de ontwikkeling van het hart. Ook spelen meerdere signaal-transductie paden een belangrijke rol. Verschillende leden van de signaal-transductie subfamilie van ‘bone morphogenetic proteins’ (Bmp)s werken nauw samen met T-box transcriptiefactoren. De Bmp subfamilie kent meer dan 20 leden, van welke in dit proefschrift vooral Bmp2 en de intracellulaire moleculen Smad1, 5, 8 en 4, die uiteindelijk de celkern in migreren na extracellulaire Bmp activatie, in detail worden besproken. Nppa, het gen dat codeert voor ‘atriale natriuretische factor’ (ANF), waarvan we in onze studie in hoofdstuk 2 de regulatoire DNA sequenties beschrijven, is een typisch voorbeeld van een doelwit-gen.

In **Hoofdstuk 1** bespreken we recent verkregen inzichten in enige vormbepalende processen die een centrale rol spelen tijdens de vroege hartontwikkeling. Daarnaast gaan we in op de implicaties van deze nieuwe inzichten voor ons denken over het ontstaan van aangeboren hartafwijkingen. We beginnen met een bespreking van ons huidige begrip van patroonvorming van de in eerste instantie rechte hartbuis langs de verschillende assen van het lichaam (cranio-caudaal (kop-staart), dorso-ventraal (rug-buik), links-rechts). Patroonvorming maakt dat de cellen in de hartbuis ‘weten’ waar ze zich bevinden, en zich als zodanig gedragen en zich verder ontwikkelen. We leggen uit wat hierbij mis kan gaan en geven inzicht in verschillende relatief veel voorkomende aangeboren afwijkingen, waaronder isomerie van de atria. Vervolgens gaan we in op de huidige kennis over de ontstaanswijze van de atriale en ventriculaire kamers, en de moleculaire regel-netwerken die hieraan ten grondslag liggen. In vrijwel alle tekstboeken over hartontwikkeling wordt nog altijd het segmentmodel beschreven. In dit model wordt er ten onrechte van uit gegaan dat in de initiële hartbuis alle voorlopercellen van de bouwstenen van het volwassen hart aanwezig zijn. Deze voorlopercellen zouden bovendien lineair gerangschikt zijn als opeenvolgende over de gehele diameter verlopende voorloперsegmenten. Hier bespreken we het ‘ballooning model van kamer ontwikkeling’. Dit is het eerste model dat inzichtelijk maakt hoe een initieel enkelvoudige buis, met een bloedstroom door een enkele holte, zich kan ontwikkelen tot een vierkamerig hart waarin de long- en de lichaams-circulatie gescheiden functioneren. Als laatste bespreken we recente inzichten in de afkomst van die cellen die worden toegevoegd nadat de embryonale hartbuis is gevormd, en over de uiteindelijke plaats en functie van deze cellen in het volwassen hart.

In **Hoofdstuk 2** beschrijven we ons onderzoek naar de regulatoire DNA sequenties die de expressie van het gen Nppa reguleren in het zich ontwikkelende en in het zieke hart. Voor de geboorte komt Nppa specifiek tot expressie in de zich vormende atriale en ventriculaire kamers. Na de geboorte wordt het gen uitgezet in de ventrikels, waar het weer wordt aangezet bij het optreden van hypertrofie (verdikking van de hartspier) en bij hartfalen. Algemeen
wordt aangenomen dat deze re-activering plaats vindt als onderdeel van de re-activering van een groter embryonaal genprogramma. Om inzicht te krijgen in de lokatie van de regulatoire sequenties van Nppa, en zo op termijn in de eiwitten die hieraan binden en dit gen reguleren, hebben we verschillende transgene muizen gegenereerd. Alle muizen hebben een ander transgeen (lichaamsvreemd) DNA construct geïntegreerd in hun erfelijk materiaal (in hun genoom), waarin steeds een ander stuk regel-DNA is gekloneerd vanuit de natuurlijke Nppa locus, dat een ‘reportergen’ aanstuurt. Een reportergen is een gen (bijvoorbeeld groen fluorescerend proteïne (GFP), LacZ of Cre) waarvan het expressiepatroon makkelijk gedetecteerd kan worden in het transgene dier. Vervolgens hebben we hartfalen geïnduceerd in de reportermuizen door ze te kruisen met een andere transgene muis die een uit gist afkomstige transcriptiefactor (Gal4) tot expressie brengt in alle hartspiercellen. Hierdoor ontstaat drie weken na de geboorte aspecifieke cardiomyopathie (ziekte van de hartspier) in alle mannetjes van deze lijn. In enkele van de reportermuizen hebben we bovendien hypertrofie geïnduceerd door ‘aortic banding’. Dit is een operatie waarbij om de aorta, net nadat deze ontspringt uit het linker ventrikel, een bandje wordt gelegd die de aorta vernauwt. Hierdoor ontstaat een druk-overbelasting, waardoor een type hypertrofie wordt geïnduceerd die ook bij bijvoorbeeld langdurige hypertensie (hoge bloeddruk) wordt gezien. Vergelijking van de expressiepatronen van de reportergenen in de diverse transgene muizen, zowel tijdens het embryonale stadium als in het zieke hart, met het expressiepatroon van endogeen (lichaamseigen) Nppa, gaf vervolgens inzicht in de potentie van de diverse regulatoire DNA sequenties. In voorafgaande studies werden alleen korte regulatoire sequenties getest. Uit deze studies werd geconcludeerd dat een dicht bij het gen liggend (proximaal) DNA fragment van 0.7 kilobasenparen alle regulatoire modules bevat die nodig zijn voor een correcte expressie voor de geboorte. De sequenties die Nppa expressie in het zieke hart aansturen werden in eerdere studies nooit gelokaliseerd. In onze studie onderzoeken we reportermuizen die korte regulatoire fragmenten op een willekeurige plek in het genoom hebben geïntegreerd, met een muis die een enkele kopie van een kort fragment heeft geïntegreerd op een welomschreven plek in het genoom (knockin muis), met muizen met een groot regulatoir fragment (tot 200 kilobasenparen in ‘bacterial artificial chromosomes’ (BACs)), en komen zo tot heel andere conclusies. We laten zien dat de proximale fragmenten reguleren elementen missen voor correcte ventriculaire activiteit tijdens de ontwikkeling. Vervolgens laten we zien dat voor correcte ventriculaire expressie voor de geboorte minimaal twee extra, op grote afstand (distaal) van het eigenlijke gen gelokaliseerde, activerende regulatoire sequenties nodig zijn. Eén van deze verzorgt de correcte patroon, de andere sequentie verzorgt de correcte embryonale activiteit. Ook de regulatoire sequenties die Nppa expressie in tijden van cardiale ziekte (stress respons) reguleren blijken distaal gelegen. Mogelijk overlappen deze, of werken samen, met de sequenties die het embryonale expressiepatroon reguleren. Echter, ze blijken onafhankelijk te functioneren van de activerende sequenties die de ventriculaire activiteit voor de geboorte reguleren. Om die reden concluderen we dat de ventriculaire Nppa activiteit voor de geboorte gereguleerd wordt door andere regulatoire sequenties en dus door andere transcriptie-paden dan de re-activatie van Nppa in tijden van cardiale stress. Deze nieuwe
informatie impliceert dat alle voorgaande data over genregulatie, verkregen uit onderzoek aan transgene muismodellen met enkel korte regulatoire Nppa fragmenten, opnieuw moeten worden geïnterpreteerd. Ook impliceert dit inzicht dat data over genregulatie die zijn verkregen uit embryonale muismodellen niet automatisch mogen worden geëxtrapoleerd naar situaties waarbij cardiale stress in het spel is, en vice versa.

**Hoofdstuk 3** bevat een studie, die al bestaande kennis verdiept over de moleculaire netwerken die de gelokaliseerde formatie van de atriale en ventriculaire kamers, versus het tussenliggende atrioventriculaire kanaal, reguleren. Voorgaande studies hadden al aangetoond dat de T-box transcriptiefactoren Tbx2, Tbx3 en Tbx20, in nauwe samenwerking met het Bmp/Smad signaal transductie pad, een centrale rol spelen in de ontwikkeling van de kamers (Tbx20) en het atrioventriculaire kanaal (Tbx2, Tbx3 en Bmp/Smad). Echter, hun onderlinge hiërarchie en precieze samenspel in tijd en plaats waren nog onduidelijk. Knock-out muizen zijn muizen waarin de werking van één specifiek gen, van een totaal van meer dan 20.000 genen, is kapot gemaakt. Door een knock-out muis te onderzoeken, kan vervolgens de functie van het specifieke gen worden afgeleid. In de hier beschreven studie hebben we muizen onderzocht waarin of alleen Tbx2, of alleen Tbx3, of zowel Tbx2 als Tbx3 was uitgeschakeld. Hieruit komt naar voren dat de transcriptiefactoren Tbx2 en het genetisch zeer verwante Tbx3 onafhankelijk van elkaar voldoende zijn om zowel de ontwikkeling van de spiercellen van het atrioventriculaire kanaal te reguleren, als de formatie van de atrioventriculaire kussens (voorlopers van kleppen en tussenschotten) te initiëren, en bovendien de kamerontwikkeling in het atrioventriculaire kanaal te onderdrukken. Kortom, beide genen zijn in het atrioventriculaire kanaal ‘redundant’ actief, wat wil zeggen dat ze essentiële functies van elkaar kunnen overnemen, mocht het andere gen niet werkzaam zijn. Vervolgens hebben we verschillende Tbx2-replicemuisen bestudeerd en de data hiervan gecombineerd met data uit experimenten waarbij we het gedrag van diezelfde reporsterconstructen in vitro (buiten het lichaam) testen door de constructen tijdelijk te introduceren in cellen. Het voordeel van deze laatste proefopzet is dat een laboratoriumsituatie wordt gecreëerd, waarbij heel specifiek de gevoeligheid van elk construct voor allerlei factoren kan worden getest. Aldoende hebben we een Bmp/Smad-afhankelijk regulatoro DNA-fragment geïdentificeerd, dat zowel de atrioventriculaire kanaal-specifieke expressie van Tbx2, als de Tbx20-afhankelijke repressie van Tbx2 in de kamers reguleert. Vervolgens laten we zien dat Tbx20 een dubbele rol speelt in de compartimentalisatie van het hart. Ten eerste blijkt, in tegenstelling tot wat tot nu toe werd verondersteld, dat Tbx20 kamerontwikkeling stimuleert onafhankelijk van Tbx2. Tegelijkertijd onderdrukt Tbx20 de activatie van Tbx2 in de zich onwikkellende kamers, waardoor Tbx2 expressie tot het atrioventriculaire kanaal wordt beperkt. Daarbij onderdrukt Tbx20 Tbx2 niet rechtstreeks door te binden aan de regulatoire DNA sequenties van Tbx2, maar indirect door te binden aan geactiveerd Smad1 en Smad5, waardoor deze eiwitten gescheiden blijven van Smad4. In het atrioventriculaire kanaal wordt Tbx2 geactiveerd door Bmp2 op een Smad-afhankelijke manier. Als resultaat van de Bmp2-binding moet, als onderdeel van de geactiveerde intracellulaire signalascade, Smad4 als cofactor eerst binden aan Smad1 of Smad5 voordat dit gehele Smad-complex vanuit het
cytoplasma de celkern in kan migreren om de transcriptie van Tbx2 te activeren. Daardoor onderdrukt Tbx20 op een indirecte manier (namelijk, door te voorkómen dat er geactiveerde Smad-complexen ontstaan die de celkern in migreren om hier Tbx2 te activeren) de Bmp/Smad-gemiedeerde activatie van het doelwit-gen Tbx2 zeer effectief. Samenvattend suggereren onze data dat tegengestelde regulatie van Bmp-signalering door Tbx20 en Tbx2 aan de basis ligt van de specificatie van respectievelijk de kamers en het atrioventriculaire kanaal.

In Hoofdstuk 4 beschrijven we het genereren van een nieuwe reportermuis, welke in zijn genoom een construct draagt waarin een groen fluorescerend eiwit (GFP) wordt aangestuurd door 160 kilobasenparen van de regulatoire sequenties van de T-box transcriptiefactor Tbx3. Terwijl endogeen Tbx3 tot expressie komt in de verschillende componenten van het cardiale geleidingssysteem, namelijk in de sinusknoop, de atrioventriculaire knoop, de atrioventriculaire (His) bundel en de bundeltakken, laten we zien dat GFP heel specifiek tot expressie komt alleen in de zich ontwikkelende atrioventriculaire knoop en niet in de andere componenten van het geleidingssysteem. Deze vinding toont aan dat de verschillende componenten van het geleidingssysteem worden gereguleerd door verschillende regulatoire DNA sequenties, en dus door verschillende transcriptionele paden. Vervolgens hebben we deze reportermuizen, die gekenmerkt worden door een fel groen fluorescerende atrioventriculaire knoop, gebruikt om sterk gezuiverde atrioventriculaire knoopcel-populaties te verkrijgen op twee verschillende stadia van ontwikkeling (embryonaal E10.5 en foetaal E17.5). Vervolgens haben we het genexpressieprofiel (de specifieke hoeveelheid RNA van alle individuele genen) bepaald van deze atrioventriculaire knoopcellen, en dit profiel vervolgens vergeleken met het genexpressieprofiel van kamerspiercellen van dezelfde leeftijd (die we hebben gezuiverd door gebruik te maken van de transgene muizen beschreven in hoofdstuk 2). Hiertoe hebben we gebruik gemaakt van microarrays, een moderne techniek die het mogelijk maakt de expressie van meer dan 20.000 genen simultaan te bepalen. Uit de hieruit gegeneerette data hebben we een volledige lijst geconstrueerd van genen die geassocieerd zijn met elektrische impulsvoortgeleiding, en die specifiek in de atrioventriculaire knoopspiercellen, dan wel in de kamerspiercellen tot expressie kwamen. Vele van deze zijn ionkanalen die niet eerder werden geassocieerd met knoop- of werk-spiecercellen, terwijl verschillende zelfs niet eerder werden geassocieerd met het hart. Bovendien komt uit deze data naar voren dat hoewel de fenotypen van de kamerspiercellen en van de atrioventriculaire knoopspiercellen gedurende de ontwikkeling divergeren, binnen elk subtype de aanwezigheid van de karakteristieke functionele genklassen grotendeels behouden blijft. Eén van de functionele genrepertoires die we hebben geïdentificeerd in de atrioventriculaire knoopcellen, bestaat uit genen die voorheen alleen werden geassocieerd met de ontwikkeling van zenuwen. Dit suggereert dat knoopspiercel-ontwikkeling en zenuw-ontwikkeling vele kenmerken delen. De informatie die uit deze microarray naar voren komt representeert het eerste genexpressieprofiel van de zich ontwikkelende atrioventriculaire knoop op genoombreed niveau en geeft ons waardevolle informatie over zijn moleculaire identiteit.
Dankwoord

‘Jongens waren we - maar aardige jongens.’ Eén van de mooiste zinnen uit de Nederlandse literatuur, waar ik vaak en lang met vrienden over gesproken heb tijdens mijn studenten- en co-schaptijd. Een boek schrijven wilden we. Daarbij ging het om de eerste en de laatste zin. Als die één keer op papier zouden staan – en er waren ideeën te over – zou het invullen van die 300 bladzijden er tussenin een koud kunstje zijn, waarna groote oplagen en beroemdhed in het verschiet zouden liggen. Nou, het boek is er bij deze gekomen. Het blijkt er echter geen van Literatuur, maar van Wetenschap. En de invulling tussen de eerste en laatste regel had meer voeten in de aarde dan toentertijd verwacht. Ook het romantisch idee van geïnspireerd schrijver (of wetenschapper) eenzaam achter zijn typemachine op zolder bleek onjuist. Dit proefschrift is tot stand gekomen door noeste arbeid en bovenal door fantastisch teamwork binnen de afdeling Anatomie & Embryologie (AEL), de overige afdelingen binnen het HartFaalCentrum, en enige onderzoeksgroepen daarbuiten. Vanaf deze plaats dank ik iedereen die op enige wijze aan dit proefschrift heeft bijgedragen, waarbij ik natuurlijk enkele mensen in het bijzonder wil noemen.

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Thomas