Exploring the neural basis of touch through selective and stable genetic tagging in the chick somatosensory system

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IV. List of abbreviations

µ Micro
%
°C Degree Celsius
AHP Afterhyperpolarization
Avil Advillin
BDNF Brain-derived neurotrophic factor
BMP Bone morphogenetic protein
bp Base pair
BSA Bovine serum albumin fraction V
Btx Bungarotoxin
CAG CMV early enhancer / chicken β-actin
cDNA Complementary DNA
chr Chromosome
cKO Conditional knockout
CNS Central nervous system
DEPC Diethylpyrocarbonate
DF Dorsal funiculus
DIG Digoxigenin
DMSO Dimethyl sulfoxide
DNA Deoxyribonucleic acid
DRG Dorsal root ganglion
E Embryonic day
ECR Evolutionary conserved non-coding region
EDTA Ethylenediaminetetraacetic acid
eGFP Enhanced GFP
EGTA Ethyleneglycoltetraacetic acid
etc Et cetera
g Gram
IV. List of abbreviations

galGal  Gallus gallus
GDNF   Glia-derived neurotrophic factor
GFP    Green fluorescent protein
GPI    Glycosyl-phosphatidyl-inositol
GRE    Gene regulatory element
h      Hour
HPD    Half peak duration
HH     Hamburger-Hamilton
IA     Intermediately adapting
IB4    Isolectin B4
Isla1  Islet1
kb     Kilo base pair
KO     Knockout
l      Liter
LF     Lateral funiculus
LT     Lateral tract
LTMR   Low-threshold mechanoreceptor
m      Milli
mm     Mus musculus
M      Molar
mGFP   Myristoylated GFP
min    Minute
MN     Motor neuron
Mrgpr  Mas-related G protein-coupled receptor
mRNA   Messenger RNA
NCC    Neural crest cell
NeuN   Neuronal nuclei
NF     Neurofilament
NGF    Nerve growth factor
NGL    NetrinG ligand
Ngn    Neurogenin
NT-3   Neurotrophin 3
Ntng   NetrinG
## IV. List of abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>P</td>
<td>Postnatal day</td>
</tr>
<tr>
<td>Phal</td>
<td>Phalloidin</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffered saline</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
</tr>
<tr>
<td>PN</td>
<td>Peripheral nerve</td>
</tr>
<tr>
<td>RA</td>
<td>Rapidly adapting</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>SA</td>
<td>Slowly adapting</td>
</tr>
<tr>
<td>SCG</td>
<td>Sympathetic chain ganglion</td>
</tr>
<tr>
<td>SDS</td>
<td>Sodium dodecyl sulfate</td>
</tr>
<tr>
<td>SEM</td>
<td>Standard error of the mean</td>
</tr>
<tr>
<td>Sox10</td>
<td>SRY box 10</td>
</tr>
<tr>
<td>STEVE</td>
<td>Stable expression vector</td>
</tr>
<tr>
<td>TF</td>
<td>Transcription factor</td>
</tr>
<tr>
<td>TFBS</td>
<td>Transcription factor binding site</td>
</tr>
<tr>
<td>TH</td>
<td>Tyrosine hydroxylase</td>
</tr>
<tr>
<td>Trk</td>
<td>Tropomyosin receptor kinase</td>
</tr>
<tr>
<td>Trp</td>
<td>Transient receptor potential cation channel</td>
</tr>
<tr>
<td>w/v</td>
<td>Weight per volume</td>
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1. Introduction

1.1. Somatosensation

Being exposed to myriad forces and environmental signals, higher organisms adapted by acquiring mechanisms to detect external stimuli, to process the information and to react accordingly. Force sensing is essential for nearly all behaviors that range from avoiding bodily harm to social exchange and is therefore fundamental for development and survival (Lumpkin et al., 2010). Many types of specialized receptors transduce light, heat, mechanical or chemical stimuli into a change in the cell’s membrane potential, the common language of neurons (Squire et al., 2008). Somatosensory neurons detect the information about noxious stimuli, touch, temperature and proprioception in skin, viscera and muscle and bridge the distance between peripheral transduction and central processing in the central nervous system (CNS) (Lewin and Moshourab, 2004; Bourane et al., 2009; Lumpkin et al., 2010). These pseudo-unipolar neurons cluster in the dorsal root ganglia (DRGs) and extend one axon collateral branch to the periphery, whereas the other collateral branch penetrates the spinal cord. The central axon collaterals form, depending on the type of stimulus they relay, synapses upon second order neurons in the dorsal horn, in the intermediate spinal cord or directly connect with motor neurons in the ventral horn (Figure 1.1) (Li et al., 2011). The CNS then constructs a topographical representation of the body, compares the input with further sensory signals and past events and processes the information, enabling animals to adapt their behavior to the changing environment (Squire et al., 2008).

All species rely on touch-dependent behaviors ranging from the simplest activities in daily life to extremely sophisticated tasks, like highly precise object manipulation or braille reading (Luo et al., 2009; Lumpkin et al., 2010). Different types of functionally specialized mechanoreceptors along with their corresponding receptor end organs in the skin encode the different qualities of touch and selectively respond to vibration, static indentation or stretch (Heidenreich et al., 2011; Li et al., 2011). The receptor density covering the body’s surface is tightly correlated with the spatial acuity, ranging from higher frequency at the finger tips to lower frequency at the back (Squire et al.,...
The interplay of the distinct mechanoreceptive subtypes provides sensory richness to touch-related percepts and encodes complex textures with fine tactile acuity (Bourane et al., 2009; Luo et al., 2009). Additionally, nociceptors and proprioceptors provide an organism with knowledge of pain sensations for avoidance behaviors and give feedback to control and coordinate muscle action, respectively (Squire et al., 2008; Arber, 2012).

Dysregulation of sensory signaling caused by disease or injury can lead to hypersensitivity and chronic pain (Lumpkin et al., 2010). Recent studies reported of a congenital insensitivity to pain caused by gene mutations (Frenzel et al., 2012). Further groups provided evidence that shared genetic factors influence different mechanosensitive systems, such as hearing and touch (Frenzel et al., 2012). Despite more than 100 years of study and its overwhelming importance in daily life, surprisingly little is known about the cellular substrate of touch sensations, including the specification of the corresponding primary somatosensory neurons, their unique function and the logic of mechanoreceptive circuit organization (Marmigère and Ernfors, 2007; Lumpkin et al., 2010; Nilius, 2010; Li et al., 2011).
1. Introduction

1.2. Somatosensory neurogenesis

The assembly of precise neuronal circuits is crucial for the function of the nervous system and depends on the specification of well-defined neuronal subpopulations that arise during embryogenesis (Kandel et al., 2000). Neurulation is initiated by the induction of the neural plate from the dorsal ectoderm of the gastrula-stage embryo, whereby the most anterior region of the neural plate forms the forebrain, progressively more posterior regions give rise to midbrain, hindbrain and spinal cord structures (Figure 1.2 a) (Tanabe and Jessell, 1996). The neural plate is subsequently converted into a tube by the shaping and folding of the neuroepithelium, involving coordinated changes in cell shape, cell division, cell migration and cell-cell interaction (Figure 1.2 b) (Squire et al., 2008; Greene and Copp, 2009). Finally, the inner neural tube and the outer surface ectoderm are created by adhesion, fusion and remodeling of the midline (Figure 1.2 c) (Copp et al., 2003). The patterning of the neuraxis is mediated by factors released from the axial and paraxial mesoderm (Tanabe and Jessell, 1996; Lumsden and Krumlauf, 1996).

These secreted factors induce the localized expression of cell-intrinsically acting factors along the rostrocaudal and dorsoventral axis of the neural tube, which, in turn, are involved in the generation of a large diversity of neuronal cell types (Muhr et al., 1999; Le Dréau and Martí, 2012). The dorsal cell fate determination depends on a cascade of secreted proteins, which are initiated by cells of the epidermal ectoderm and propagated by roof plate cells within the neural tube. The main instructive cues are members of the Transforming growth factor β/Activin/Bone morphogenetic protein (BMP) family, complemented by additional signals involving the Wnt and Fibroblast growth factor families, which also contribute to the dorsal neuronal identity (Liem et al., 1997; Lee and Jessell, 1999; Helms and Johnson, 2003; Le Dréau and Martí, 2012). The ventral neuronal patterning of the CNS is induced by signaling of the notochord and later on by the floor plate through secretion of the glycoprotein Sonic hedgehog, which has been shown to direct different cell fates, such as motor neurons and ventral interneurons, at different concentration thresholds (Chiang et al., 1996; Briscoe et al., 2000). Selective cross-repressive interactions convert the dorsoventral
signaling activity into discrete progenitor domains (Sander et al., 2000; Liem et al., 2000; Jessell, 2000).

Concomitant with the fusion of the neural folds during neurulation, neural crest cells (NCCs) subsequently emerge from the most dorsal portion of the neural tube. These multipotent, SRY box 10 (Sox10)-positive progenitor cells give rise to a wide variety of cell types and form most of the peripheral nervous system, including dorsal root, sympathetic, parasympathetic and enteric ganglia. Furthermore, NCCs generate non-neuronal cells like glia, Schwann cells, adrenal cells and melanocytes (Bronner-Fraser and Fraser, 1988; Frank and Sanes, 1991; Gilbert, 2000; Kim et al., 2003). NCC progenitor cell fate arises prior to neurulation within a band of cells at the border between the neural plate and the non-neuronal ectoderm (Le Douarin and Kalcheim, 1999; Squire et al., 2008). Premigratory NCC development is induced by signaling factors, including BMP and WNT, whereas a variety of further factors can bias the cells toward certain lineages (Bronner-Fraser, 2004; Huang and Saint-Jeannet, 2004;
NCCs undergo a transition from epithelial to migratory mesenchymal cells with changes in their adhesive properties (Sauka-Spengler and Bronner-Fraser, 2008; Squire et al., 2008). Downregulation of cell adhesion molecules like N-cadherin and cadherin 6 are thought to drive the transition from epithelial to motile mesenchymal characteristics (Nakagawa and Takeichi, 1998; Pla et al., 2001). During this process, the basal lamina around the neural tube dissolves, NCCs delaminate and directionally emigrate, a process involving homotypic cell-cell interactions (Kandel, 2000; Carmona-Fontaine et al., 2008). Many NCCs appear to be multipotent both before and after emigration, and their fate is to large degree determined in response to extracellular signals (Hari et al., 2012). However, several lines of evidence suggest the existence of subpopulations of lineage-restricted premigratory NCCs (George et al., 2007; Lefcort and George, 2007). NCCs migrate along defined pathways in chain-like structures towards their respective peripheral targets between embryonic day (E) 8 and E10 in the mouse and between Hamburger-Hamilton stage 11 and 21 in the chick (Hamburger and Hamilton, 1951; Serbedzija et al., 1990; Frank and Sanes, 1991; Kasemeier-Kulesa et al., 2005; Krispin et al., 2010). Attractive and repulsive environmental adhesion molecules and secreted guidance cues influence NCC migration before they differentiate into their target-appropriate cell types (Ruhrberg and Schwarz, 2010; Theveneau and Mayor, 2012). Currently, little is known about how NCCs stop their migration after reaching their appropriate destination (Squire et al., 2008; Theveneau and Mayor, 2012).

Neurogenesis occurs in three successive waves of NCC migration between E9.5 and E11 in the mouse, in which the Wnt/β-catenin pathway and the BMP signaling play an instructive role concerning the specification of the sensory lineage (Hari et al., 2002; Lee HY et al., 2004; Bronner-Fraser, 2004; Kléber et al., 2005; Marmigère and Ernfors, 2007). After their delamination a subpopulation of NCCs migrates along a ventral pathway and coalesces into dorsal root ganglia (DRGs) adjacent to the neural tube at regular intervals (Lallemand and Ernfors, 2012). The first wave of neurogenesis is initiated during early NCC migration by the expression of the transcription factor (TF) Neurogenin 2 (Ngn2) that directs NCC differentiation mainly into mechanoreceptive and proprioceptive sensory neurons, which localize in the ventrolateral part of the DRG (Figure 1.3 a) (Ma et al., 1999; Perez et al., 1999). These
Ngn2⁺ cells show limited cell division with three neurons per NCC and constitute around four percent of the adult DRG neurons (Frank and Sanes, 1991; Anderson, 2000).

With around 91%, the major proportion of the DRG neurons is generated in the second wave of sensory neurogenesis due to a high cell division rate that produces an average of 36 neurons per NCC (Figure 1.3 b) (Frank and Sanes, 1991; Marmigère and Ernfors, 2007). High levels of the TF Neurogenin 1 (Ngn1) drive the postmigratory differentiation of the multipotent Sox10⁺ NCCs towards small nociceptive neurons located in the dorsomedial region of the DRG, as well as towards large-size mechanoreceptive and proprioceptive neurons (Ma et al., 1999; Kim et al., 2003). Furthermore, in a third wave of neurogenesis, boundary cap cells, a specialized population of postmigratory NCCs contributing to up to five percent of total DRG neurons, are thought to produce DRG neurons of exclusively nociceptive type.
1. Introduction

(Figure 1.3 c) (Maro et al., 2004; Lallemend and Ernfors, 2012). NCCs of the second and third waves also differentiate into peripheral glia in addition to neurons (Frank and Sanes, 1991; Maro et al., 2004). The Neurogenin TFs initiate pan-neuronal programs leading to the establishment of neuronal fate accompanied by the expression of other key TFs like Brn3a, Islet1 or Foxs1 (McEvilly et al., 1996; Perez et al., 1999; Kim et al., 2003; Montelius et al., 2007). The initiated transcriptional programs cause downregulation of multipotency factors like Sox10 and suppression of dorsal spinal cord and other non-neuronal fates, while downstream signaling cascades promote neuronal differentiation and sensory neuron subtype specification (Marmigère and Ernfors, 2007; Sun et al., 2008; Lanier et al., 2009).

1.3. Somatosensory neuron specification and connectivity

The determination of a neuronal cell fate is controlled by the interplay between environmental cues and cell-intrinsic information. Therefore, the birth of a neuron at a specific time and position determines its identity (Lallemend and Ernfors, 2012). Shortly after their accumulation in DRGs the immature NCC-derived neurons start to differentiate into various types of sensory neurons, including nociceptive and mechanosensitive neurons terminating in the skin, as well as proprioceptive neurons innervating deep structures, such as muscle spindles and golgi tendon organs (Lupkin et al., 2010). Intrinsic and local environmental signals drive the segregation into unique subtypes by transcriptional activation and repression of specific genes, encoding key cell fate determinants. The earliest known markers for sensory subtypes are the growth factor receptors Tropomyosin receptor kinase A (TrkA), TrkB, TrkC, Met and Ret which serve as receptors for the Neurotrophins (Nerve growth factor, NGF; Brain-derived neurotrophic factor, BDNF; Neurotrophin 3, NT-3), Hepatocyte growth factors and Glia-derived neurotrophic factor (GDNF) ligands produced by cells from sensory target tissue (Snider and Wright, 1996; Huang and Reichardt, 2001).
These factors and their receptors play important roles in shaping the functionality and specific connectivity of the primary somatosensory neurons by their involvement in neuron diversification, peripheral innervation, cell survival and central axon collateral targeting (Huang et al., 1999; Fünfschilling et al., 2004; Marmigère and Ernfors, 2007; Luo et al., 2009; Gascon et al., 2010). Furthermore, several differentially expressed TFs begin to be expressed in early neurogenesis, including MafA and c-Maf, as well as the Runx family members Runx1 and Runx3 (Kramer et al., 2006; Bourane et al., 2009; Wende et al., 2012). It is generally thought that these factors participate via combinatorial expression profiles in the basic establishment of sensory neuron diversification and maturation, a process that extends well into postnatal stages in mammals (Figure 1.4). Genetic programs that direct neuronal subtype specification are likely to also control connectivity. However, the regulatory mechanisms driving
both sensory subtype specification and connectivity remain incomplete and additional participating factors need to be identified (Liu and Ma, 2011; Lallemend and Ernfors, 2012).

1.3.1. Nociceptive neurons
The major part of unmyelinated nociceptive neurons is generated from the Ngn1-dependent wave of neurogenesis, followed by the upregulation of high TrkA and Runx1 expression levels at E11.5 in mice (Ma et al., 2003; Kramer et al., 2006; Chen CL et al., 2006). Runx1 is important for the emergence of Ngn1-dependent small-diameter neurons by activation of TrkA via direct promoter binding (Marmigère et al., 2006). At later embryonic stages around E12.5 - E15.5 in mice, Runx1 expression becomes more subtype-restricted and the function switches from a general nociceptive transcriptional activator to a repressor (Chen CL et al., 2006). Thus, Runx1 mediates the further diversification of immature TrkA⁺ neurons. Maintained Runx1 expression drives a TrkA⁻ non-peptidergic phenotype, whereas downregulation of Runx1 allows neurons to acquire a TrkA⁺ peptidergic phenotype (Chen CL et al., 2006; Marmigère and Ernfors, 2007).

Figure 1.5: Connectivity of nociceptive neurons

(a) Nociceptive subtypes terminate in different laminae in the spinal cord. Peptidergic C-fibers innervate laminae I and II (outer), non-peptidergic neurons project mainly into lamina II (inner), whereas lightly myelinated Aδ-nociceptors terminate in laminae I and V of the dorsal horn. 
(b) In the periphery, peptidergic C-fibers terminate as free nerve endings in the epidermis close to the dermis border and as circular nerve endings at hair follicles. Non-peptidergic peripheral projections terminate superficially in the epidermis.

(Adapted from Lallemend and Ernfors, 2012)
Peptidergic nociceptors, which serve as polymodal pain-sensing neurons responding to noxious heat and mechanical stimuli, express high levels of TrkA, Met and the neuropeptides CGRP and Substance P (Lewin and Moshourab, 2004). Their slowly conducting C-fibers terminate close to the epidermis-dermis border as free nerve endings and as circular nerve endings at hair follicles (Figure 1.5). Their central afferents project into laminae I and II (outer) in the spinal cord for pain transmission (Fitzgerald, 2005; Zylka et al., 2005; Squire et al., 2008). Activation of TrkA has been shown to induce Met, which defines the peptidergic phenotype by extinguishing Runx1 expression via cross-repressive interactions between Met and Runx1 (Kramer et al., 2006; Gascon et al., 2010). Additionally, Met and TrkA activate peptidergic expression profiles (Lumpkin and Caterina, 2007; Lallemend and Ernfors, 2012).

Non-peptidergic nociceptors are responsible for the detection of thermal pain, inflammatory pain and are involved in neuropathic pain (Chen CL et al., 2006). Their C-fiber terminals innervate the epidermis and their central projections terminate in lamina II (inner) in the spinal cord (Figure 1.5) (Fitzgerald, 2005; Zylka et al., 2005; Squire et al., 2008). Non-peptidergic neurons display a unique expression of Runx1 and Ret, which together induce the progressive extinction of TrkA around P14 in mice, while at the same time these neurons acquire isolectin B4 (IB4) binding (Molliver et al., 1997; Abdel Samad et al., 2010). During segregation of the peptidergic and non-peptidergic subtypes, Runx1 is important for inducing Ret expression and for repression of the peptidergic neuron markers CGRP and Met (Luo et al., 2009; Gascon et al., 2010). The upregulation of non-peptidergic-specific ion channels and transmembrane receptors shapes the receptive properties and the neuronal connectivity (Lumpkin and Caterina, 2007; Golden et al., 2010). Additionally, Ret, whose expression is initiated by NGF and consolidated by Ret itself, accompanies the Runx1 function in repressing TrkA during the segregation of the lineages (Luo et al., 2007). At approximately E16.5 in mice further nociceptive subtype specification occurs by the expression of different Mas-related G protein-coupled receptors (Mrgprs) whose functions are thought to underlie the acquisition of distinct pain sensitivities (Liu et al., 2008; Lallemend and Ernfors, 2012).
It has been shown, that Runx1 is also necessary for the specification of DRG neurons expressing the Transient Receptor Potential channel family member TrpM8 (Chen CL et al., 2006). These TrpM8+, but CGRP– and IB4– unmyelinated or lightly myelinated neurons probably arise from early TrkA+/Runx1+ progenitors and mediate innocuous and painful sensations to cold and cooling compounds (Peier et al., 2002; Takashima et al., 2010). New data further revealed an early TrkA lineage of Aδ-nociceptors, generated at E10.5 by a Ngn1-independent neurogenesis terminating in laminae I and V of the dorsal horn (Lallemend and Ernfors, 2012). This DRG neuron subtype expresses CGRP and the myelination marker Neurofilament 200 (NF200), whereas no expression of Runx1, Ret or Met could be detected (Gascon et al., 2010; Bachy et al., 2011). In summary, based on current literature, at least two bursts of birth of TrkA+ neurons exist in sensory neurogenesis (Figure 1.4).

1.3.2. Proprioceptive neurons

Large-size proprioceptive neurons play a prominent role in coordinated muscle control by detecting changes in muscle length (Arber, 2012). Type Ia heavily myelinated Aα-fibers innervate muscle spindles, whereas the central afferents project to the intermediate and ventral spinal cord where they directly synapse with α-motor neurons to establish monosynaptic reflex circuits (Chen et al., 2003). Group Ib Aα-afferents which innervate golgi tendon organs project mainly to lamina VI in the intermediate spinal cord. Proprioceptive Aβ-fibers of group II DRG neurons also terminate at muscle spindles and, like type Ia afferents, concentrate their central afferents in the intermediate and ventrolateral gray matter (Figure 1.6) (Scott, 1992; Riddell and Hadian, 2000).

The progenitors that give rise to proprioceptive neurons arise from the first and second waves of neurogenesis in the DRG (Ma et al., 1999). Early in sensory DRG neuron diversification, two TrkC+ populations can be observed until E12.5 in mice: one population expresses TrkC in combination with TrkB, whereas the other one expresses TrkC alone. At approximately E10.5, expression of Runx3 is initiated leading to the repression of TrkB and Shox2 (Kramer et al., 2006; Abdo et al., 2011; Scott et al., 2011). This direct repression can be explained by the dependence of a
TrkB gene silencer on the Runx protein complex and on NT-3 signaling (Inoue et al., 2007).

Figure 1.6: Connectivity of proprioceptive neurons
(a) Type Ia proprioceptive neurons terminate in the ventrolateral spinal cord and directly connect motor neurons (IX). Type Ib afferents project to the intermediate zone (mainly lamina VI), whereas type II neurons occupy both intermediate and ventral domains of the spinal cord.
(b) Type Ia Aα-fibers and type II Aβ-fibers innervate muscle spindles. Type Ib Aα-afferents form endings at Golgi tendon organs.

(Adapted from Lallemend and Ernfors, 2012)

The initiation of Runx3 expression in immature proprioceptive neurons occurs before peripheral target innervation and implies the regulation of determinants in the local DRG environment for proprioceptive cell fate (Lallemend and Ernfors, 2012). Additionally, Runx3 maintains TrkC expression via cross-activating loops between Runx3 and TrkC to consolidate a proprioceptive phenotype between E11 and E13.5 (Figure 1.4) (Kramer et al., 2006; Nakamura et al., 2008). The TrkC ligand NT-3, which is expressed in early muscle mass and later on in muscle spindles, has been shown to induce the ETS family TF ER81, an important regulator of central connectivity (Patel et al., 2003). Together, Runx3 and TrkC are the key factors in molecular specification and functional circuit formation of proprioceptive neurons (Inoue et al., 2002; Levanon et al., 2002; Chen Al et al., 2006). At later embryonic stages, additional general proprioceptive markers like Parvalbumin begin to be expressed, but no genes exclusively expressed by type Ia, Ib or II neurons have been identified so far (Chen et al., 2003).
1.3.3. Mechanoreceptive neurons

Mechanoreceptive neurons comprise various myelinated and unmyelinated types, terminating throughout laminae II (inner) and V in the dorsal horn of the spinal cord and innervating their corresponding peripheral end organs in the hairy and glabrous skin (Figure 1.7) (Brown, 1982; Squire et al., 2008).

Figure 1.7: Connectivity of mechanoreceptive neurons

(a) Mechanoreceptive central afferents project into deeper laminae of the spinal cord. Rapidly adapting (RA) and slowly adapting (SA) low-threshold mechanoreceptors (LTMRs) terminate throughout laminae III-V, Aδ-LTMRs mainly connect with interneurons from lamina III and unmyelinated C-afferents project into lamina II (inner).

(b) Pacinian corpuscles and Merkel cells present in both, glabrous and hairy skin, are innervated by RA-afferents and SA-afferents, respectively. Meissner corpuscles exclusively found in glabrous skin and hair follicles are also innervated by RA Aδ-LTMRs, whereas SA-afferents additionally terminate at Ruffini corpuscles in glabrous skin. Aδ- and C-fibers were found to form lanceolate and circular nerve endings at hair follicles.

(Adapted from Lallemend and Ernfors, 2012)

In contrast to proprioceptors and nociceptors, little is known about the molecular mechanisms governing the early differentiation and later specification into mechanosensitive subtypes (Lecoin et al., 2010). Mechanoreceptive neurons arise mainly from the Ngn2-dependent wave of neurogenesis (Ma et al., 1999). Soon after DRG accumulation, mechanoreceptive progenitors segregate into Ret/Gfra2 expressing (earlyRet⁺) and TrkB⁺/TrkC⁺ populations (Luo et al., 2009; Honma et al., 2010). This lineage diversifies further into Ret⁺/MafA⁺, Ret⁺/MafA⁺/TrkB⁺ and Ret⁺/MafA⁺/TrkC⁺ expressing neurons, for which Ret signaling appears to be crucial (Bourane et al., 2009; Lecoin et al., 2010). The more broadly expressed c-Maf controls MafA expression in these cells and further regulates the expression of ion channels.
important for mechanoreceptor functions (Wende et al., 2012). MafA, which is expressed in all of the three earlyRet+ mechanoreceptive subtypes, shows the ability to repress TrkB (Ma, 2009). TrkB expression in one of the subtypes can only be maintained by additional expression of Shox2 (Abdo et al., 2011). The regulatory mechanism of the differentiation into Ret+/MafA+ or Ret+/MafA+/TrkC+ mechanoreceptors still remains unknown (Lallemend and Ernfors, 2012). The earlyRet+ neurons have been shown to predominantly label rapidly adapting low-threshold mechanoreceptors (LTMRs) innervating Meissner corpuscles, Pacinian corpuscles and hair follicles (Ma, 2009). Ret, MafA and c-Maf mutant mice all show dramatic defects in rapidly adapting mechanoreceptors function, as well as their central and peripheral connectivity (Table 1.1) (Bourane et al., 2009; Luo et al., 2009; Honma et al., 2010; Wende et al., 2012). Whether the different earlyRet+ subtypes are restricted to different mechanoreceptive end organs demands further investigation. Therefore, additional participating factors, like ion channels responding to mechanical stimuli, still need to be identified (Ma, 2009). One putative candidate could be the potassium channel Kcnq4, which is expressed in earlyRet+ neurons exclusively innervating Meissner corpuscles and hair follicles and plays an important role in controlled low-frequency vibration detection (Heidenreich et al., 2011).

The TrkB+ lineage of mechanoreceptive neurons arises from early TrkB+/TrkC+ progenitors and might include slowly adapting LTMRs innervating Merkel and Ruffini end organs and lightly myelinated Aδ-LTMRs (Abdo et al., 2011; Li et al., 2011). The co-expression of Shox2 in the progenitors promotes TrkB and represses TrkC expression and thereby directs the neurons to escape from a proprioceptive phenotype (Scott et al., 2011). The cross-activation between Shox2 and TrkB and the cross-repression between Shox2 and TrkC are suggested to be the major determinants in TrkB+ mechanoreceptive versus TrkC+ proprioceptive cell fates (Lallemend and Ernfors, 2012).

Furthermore, Li and colleagues identified a DRG neuron population representing unmyelinated C-LTMRs, that is positive for Ret, tyrosine hydroxylase (TH) and vGlut3+, but negative for the common nociceptive markers TrkA, TrpV1, Mrgps and IB4 (Li et al., 2011). These recent findings imply the existence of at least five molecularly
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unique mechanoreceptive subtypes: Ret⁺/MafA⁺, Ret⁺/MafA⁺/TrkB⁺, Ret⁺/MafA⁺/TrkC⁺, TrkB⁺ and Ret⁺/TH⁺ (Figure 1.4). However, in most cases their correlation with distinct mechanoreceptive end organs needs to be examined in more detail (Table 1.1).

Table 1.1: Correlation of mechanoreceptive subtypes with their innervated end organs

<table>
<thead>
<tr>
<th>lineage</th>
<th>method</th>
<th>innervated end organs</th>
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<tbody>
<tr>
<td></td>
<td>rapidly adapting</td>
<td>Meissner corpuscles</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacinian corpuscles</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Merkel cells (hairy)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Merkel cells (glabrous)</td>
</tr>
<tr>
<td>earlyRet</td>
<td>earlyRet (genetically)</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>Ret cKO</td>
<td>yes</td>
</tr>
<tr>
<td>Luo et al., 2009</td>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>earlyRet</td>
<td>Ret (antibody)</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>Ret cKO</td>
<td>yes</td>
</tr>
<tr>
<td>Bourane et al., 2009</td>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>earlyRet</td>
<td>Ret (genetically)</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>Ret KO</td>
<td>generally impaired innervation</td>
</tr>
<tr>
<td>Hamma et al., 2010</td>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>earlyRet</td>
<td>cMaf cKO</td>
<td>affected</td>
</tr>
<tr>
<td></td>
<td></td>
<td>affected</td>
</tr>
<tr>
<td>Wende et al., 2012</td>
<td></td>
<td>lost</td>
</tr>
<tr>
<td>earlyRet subtype</td>
<td>Kcnq4 (antibody)</td>
<td>affected</td>
</tr>
<tr>
<td></td>
<td></td>
<td>unaffected</td>
</tr>
<tr>
<td>Heidenreich et al., 2011</td>
<td></td>
<td>affected</td>
</tr>
<tr>
<td>earlyRet subtype</td>
<td>Shox2 cKO</td>
<td>affected</td>
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<td></td>
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<td>unaffected</td>
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<tr>
<td>Abdø et al., 2011</td>
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<td>affected</td>
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</tbody>
</table>

The current knowledge regarding the process of sensory neuron diversification is still rudimentary. The discovery of somatosensory subtype-specific direct cell lineage tracings, co-activators, co-repressors and further downstream substrates would help to understand the definite hierarchical organization, sensory cell fate determining mechanisms or the precise connectivity of somatosensory neuronal circuits (Marmigère and Ernfors, 2007; Lallemend and Ernfors, 2012).
1.4. **Low-threshold mechanoreceptor characteristics**

Low-threshold mechanoreceptors (LTMRs) sense a range of innocuous mechanical stimuli and are subdivided according to their temporal response properties and their receptive field size. LTMRs differ in their conduction velocities which are affected by the nerve diameter, the myelination and the internal axon resistance. Accordingly, rapidly adapting (RA) and slowly adapting (SA) types can be distinguished (Lewin and Moshourab, 2004; Lumpkin and Caterina, 2007; Ma, 2009). LTMRs possessing a small receptive field size in the skin are referred to as type I, while LTMRs with a large receptive field are termed type II (Woolfe et al., 2008). In mammals, RA type I and RA type II Aβ-mechanoreceptors innervate Meissner corpuscles and Pacinian corpuscles, respectively, that dynamically respond to low and high-frequency vibration (Lewin and Moshourab, 2004; Heidenreich et al., 2011). Hair follicles are innervated by heavily myelinated RA Aβ-LTMRs, lightly myelinated Aδ-LTMRs (D-hair receptors) and unmyelinated C-LTMRs, all of which are stimulated by hair deflection (Lewin and Moshourab, 2004; Lumpkin et al., 2010). The peripheral collaterals of SA type I and SA type II Aβ-mechanoreceptors terminate in Merkel discs and Ruffini corpuscles, respectively, that relay static responses to dermal stretch and indentation (Lewin and Moshourab, 2004; Lumpkin and Caterina, 2007; Lumpkin et al., 2010). The different large-diameter Aβ-LTMRs, as well as the medium-diameter Aδ-neurons and small-diameter C-fibers can be differentiated electrophysiologically via *in vitro* skin nerve preparations, whereby response properties of single axon fibers are recorded after variable mechanical stimulation of the skin (Figure 1.8) (Lewin and Moshourab, 2004; Heidenreich et al., 2011; Wende et al., 2012).

Through whole cell patch clamp recordings, it is further possible to discriminate isolated adult and embryonic nociceptive and mechanoreceptive sensory neurons by their characteristic action potential waveforms (Koerber et al., 1988; Djouhri et al., 1998; Fang et al., 2005). The nociceptive action potentials show a ‘hump’ on the falling phase, long half peak durations and longer afterhyperpolarization durations. In contrast, mechanoreceptors fire narrow uninflected spikes with short half peak and afterhyperpolarization durations (Lechner et al., 2009). Additionally, all sensory neurons acquire a mechanotransduction competence, the ability to transduce
mechanical stimuli into electrical signals (Lechner et al., 2009). In mechanoreceptors and proprioceptors, this mechanically-activated rapid and highly sensitive mechanosensitive current activates and inactivates within a few milliseconds and appears as soon as the receptors innervate their peripheral targets (Lechner et al., 2009). LTMRs thus possess a so-called RA-mechanosensitive current, while the majority of nociceptors exhibit intermediately adapting and SA inactivating currents (Drew et al., 2002; Hu and Lewin, 2006; Lechner et al., 2009).

It is therefore possible to classify LTMRs based on their peripheral innervation of distinct morphologically unique end organs, as well as on their characteristic physiological properties. The central afferents of LTMRs also differ from other sensory types in their termination zones in the spinal cord. RA and SA Aβ-LTMRs terminate throughout laminae III to V, Aδ-LTMRs mainly connect with interneurons from lamina III and unmyelinated C-afferents project into lamina II (inner) (Light and Perl, 1979; Li et al., 2011). In contrast, the majority of nociceptive afferents project in the most superficial laminae I and II of the dorsal horn, while proprioceptive projections terminate in laminae VI and XI of the intermediate and ventrolateral gray matter.

Figure 1.8: Electrophysiology of somatosensory neuron subtypes
(a) mechanoreceptive neurons
RA Aβ-mechanoreceptors (~10%)  SA Aβ-mechanoreceptors (~12%)  Aδ-mechanoreceptors (~6%)
(b) nociceptive neurons
Aβ-nociceptors (~12%)  C-mechano-nociceptors (~20%)  C-mechanoheat-nociceptors (~40%)

In vitro mouse skin nerve preparations reveal typical electrophysiological differences between sensory neuron subtypes and their approximate incidence of total cutaneous sensory neurons. (a) Low-threshold mechanoreceptors robustly respond to the ramp phase of the stimulated skin. RA, rapidly adapting; SA, slowly adapting. (b) Nociceptors respond primarily to the static phase of the stimulus. (Modified from Lewin and Moshourab, 2004)
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(Brown, 1982; Chen AI et al., 2006). In analogy to the fine-grained laminar and sublaminar segregation patterns of the molecularly defined nociceptor identities, the different subtypes of LTMRs could consequently also terminate in discrete laminar targets (Zylka et al., 2005; Li et al., 2011). Li and colleagues could recently show that the major hair follicle types (guard hair, zigzag hair and awl/auchene hair) are innervated by a unique and invariant combination of LTMRs (Figure 1.9). Strikingly, they could also show a topographical arrangement of the corresponding central afferents in columns, whereby the projections of the different LTMRs innervating the same or adjacent hair follicles terminate within narrow columns in the dorsal horn (Li et al., 2011). The dorsal horn could therefore combine 2000-4000 columns of LTMR units in 3D space that reflect the respective peripheral receptive fields (Li et al., 2011; Arber, 2012). A similar concept has been proposed for nociceptive withdrawal reflex modules (Ladle et al., 2007; Arber, 2012).

Molecularly, at least five unique mechanoreceptive subtypes (Ret⁺/MafA⁺, Ret⁺/MafA⁺/TrkB⁺, Ret⁺/MafA⁺/TrkC⁺, TrkB⁺, Ret⁺/TH⁺) were identified, but their correlation with distinct mechanoreceptive end organs and their central terminations in the spinal cord need to be determined in more detail (Table 1.1) (Bourane et al., 2009; Luo et al., 2009; Honma et al., 2010; Abdo et al., 2011; Heidenreich et al., 2011; Li...
et al., 2011; Wende et al., 2012). Further analysis becomes difficult, however, due to the scarcity of available gene expression profiles or specific markers that would help to directly identify the mechanoreceptive subtypes (Ma, 2009; Lallemend and Ernfors, 2012).

Surprisingly, little is known about the establishment of well-defined sensory circuits. Gene programs that direct the sensory subtype specification are likely to control connectivity (Hippenmeyer et al., 2004). It has been shown that key determinants for the establishment of sensory phenotypes like Runx1, Runx3 and Ret also have an impact on the neuron subtype-specific connectivity, presumably by influencing the expression of cell-surface receptors for axon guidance cues (Chen AI et al., 2006; Chen CL et al., 2006; Kramer et al., 2006; Marmigère and Ernfors, 2007; Bourane et al., 2009; Luo et al., 2009). Recent evidence indicates that peripheral target-derived, yet-to-be-identified retrograde signals prompt sensory afferents to project to spatially stereotyped and conserved domains in the spinal cord and hindbrain (Sürmeli et al., 2011). In these confined zones presynaptic sensory axons and postsynaptic interneuron/motor neuron dendrites overlap and form synaptic contacts presumably through the selective interaction of cell-surface signaling cues (Arber, 2012). Axon target finding results therefore from interplay between long-range signaling for modality-specific innervation and attractive and repulsive short-range forces for lamina-specific synaptic connectivity (Chen et al., 2003; Marmigère and Ernfors, 2007). Target-derived neurotrophic factors like NT-3 in muscles and NGF signaling in the dermis are important participants in establishing somatosensory circuits (Patel et al., 2000; Ernfors, 2001; Patel et al., 2003). Another factor is the short-range diffusible chemorepellent Sema3a, which mediates a dorsal discrimination of nociceptive and mechanoreceptive afferents versus ventrally projecting proprioceptive afferents (Messersmith et al., 1995; Pond et al., 2002; Marmigère and Ernfors, 2007). However, the complex and unique wiring of sensory circuits is far from being understood (Arber, 2012). The identification of additional factors and downstream substrates that execute the effects of specification and connectivity would shine a light on the regulatory mechanisms of sensory neurons and bridge the gap between their molecular identity and their precise peripheral and central circuit formation (Liu and Ma, 2011; Lallemend and Ernfors, 2012).
1.5. NetrinG-NGL interactions in neuronal circuit formation

Cell adhesion molecules at neuronal synapses regulate diverse aspects of neuronal circuit formation, including axo-dendritic contact establishment, synapse formation and maturation (Woo et al., 2009). The synaptic adhesion molecules NetrinG1 and NetrinG2 appear to be expressed by a subset of large-diameter sensory neurons in postnatal mouse DRGs and are therefore candidate factors in sensory circuit formation (Yin et al., 2002; www.brain-map.org). These receptors share a similar domain structure with the well-characterized secreted Netrin axon guidance molecule. In contrast to Netrins, NetrinGs are plasma membrane-linked via a Glycosylphosphatidyl-inositol (GPI) anchor and display isoform-specific interaction with the NetrinG ligands (NGL) 1 and NGL2, respectively, which are also known as LRRC4C and LRRC4 (Nakashiba et al., 2002; Yin et al., 2002; Niimi et al., 2007; Nishimura-Akiyoshi et al., 2007). NetrinG and NGL homologs are only found in vertebrates and are supposed to be involved in aspects of the nervous system of higher organisms with greater diversity of synapses (Woo et al., 2009). Their multiple splice variants are differentially expressed in mainly non-overlapping neuronal populations in distinct regions of the brain during embryonic and postnatal developmental stages (Nakashiba et al., 2002; Yin et al., 2002; Meerabux et al., 2005; Niimi et al., 2007; Nishimura-Akiyoshi et al., 2007).

NetrinG-NGL interactions have been implicated in the regulation of axonal outgrowth and synaptic differentiation (Nakashiba et al., 2002, Lin et al., 2003; Woo et al., 2009). Transsynaptic NetrinG-NGL adhesion promotes the formation of excitatory, but not inhibitory synapses through recruitment of pre- and postsynaptic proteins like PSD95 and NMDA (Kim et al., 2006; Biederer, 2006; Brose, 2009; Linhoff et al., 2009). A study by Nishimura-Akiyoshi and colleagues associated NetrinG-NGL interactions with lamina-specific segmentation of dendrites (Nishimura-Akiyoshi et al., 2007). Entorhinal cortex axon fibers selectively expressing NetrinG1 terminate on distal dendrites of CA1 pyramidal neurons of the hippocampus, whereas NetrinG2 expressing Schaffer collaterals from CA3 neurons mainly terminate on proximal regions of the CA1 dendrites. Accordingly, NGL1 and NGL2 localize in the distal and proximal dendritic
segments of CA1 pyramidal neurons, respectively (Figure 1.10) (Nishimura-Akiyoshi et al., 2007).

Knockout mice of NetrinG1 and NetrinG2 show a selective dispersal of their ligands. Therefore, it is believed that presynaptic NetrinGs localize and stabilize postsynaptic NGLs in specific segments of contacting dendrites through isoform-specific transneuronal adhesion for further synaptic protein recruitment and input-specific synapse formation (Nishimura-Akiyoshi et al., 2007; Woo et al., 2009; Seiradake et al., 2011). Furthermore, knockout mice revealed the importance of NetrinG-NGL interactions for auditory synaptic responses (Zhang et al., 2008). Whirlin, a direct interaction partner of NGL1, has been detected in stereocilia on the outer surface of hair cells in the cochlear system and has been linked to congenital hearing impairment (Mburu et al., 2003; Delprat et al., 2005). In humans, single nucleotide polymorphism studies associated NetrinGs with schizophrenia, bipolar disorder and a rare cause of
atypical Rett syndrome (Aoki-Suzuki et al., 2005; Borg et al., 2005; Archer et al., 2006; Nectoux et al., 2007; Eastwood and Harrison, 2008; Ohtsuki et al., 2008).

Figure 1.11: Expression analysis of NetrinG1 and NGL1 in vertebrates

In summary, NetrinG-NGL interactions are short-range cues for axonal and dendritic behavior through bidirectional signaling and have a high impact on axonal outgrowth, lamina-specific dendritic segmentation and synapse formation, whereas deregulation of NetrinGs/NGLs has been implicated in diverse brain dysfunctions (Nakashiba et al., 2002; Woo et al., 2009). Based on literature and gene expression databases, Ntng1 mRNA expression is detected in a scattered subset of large soma size sensory neurons in mouse P4 transverse DRG sections. Numbers indicate presumptive laminae. Scale bar: 100 µm.

(b) Hypothesis of selective laminar targeting of NetrinG1+ mechanoreceptive central afferent projections in laminae IV-V presumably expressing NGL1 in mammals and avians.

(Modified from www.brain-map.org)

In summary, NetrinG-NGL interactions are short-range cues for axonal and dendritic behavior through bidirectional signaling and have a high impact on axonal outgrowth, lamina-specific dendritic segmentation and synapse formation, whereas deregulation of NetrinGs/NGLs has been implicated in diverse brain dysfunctions (Nakashiba et al., 2002; Woo et al., 2009). Based on literature and gene expression data bases, Ntng1 mRNA expression is detected in a scattered subset of large soma size sensory neurons in mouse P4 and P20 DRGs (Yin et al., 2002; www.brain-map.org). Interestingly, the Ntng1 ligand NGL1 is highly expressed in the mouse P4 spinal cord and appears restricted to deeper laminae of the dorsal horn, which could correspond to laminae IV-V, while being excluded from superficial laminae. Consequently, Ntng1-NGL1 interactions might be involved in the establishment of the specific and well-defined sensory connectivity during development (Figure 1.11).
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1.6. Identification of cis-regulatory elements

Precise control of embryonic development and maintenance of the organism is mediated by a complex interaction network between a limited amount of around 20,000 genes and their products (Alberts et al., 2008; Uchikawa, 2008). The identification of gene regulatory elements (GREs) is thus a first step to decipher the gene control machinery (Maher, 2012). Additionally, lineage-specific GREs can be used as genetic tools for tagging and manipulation of discrete cell types, like sensory neuron subtypes, to study their development and function (Echelard et al., 1994; Marquardt et al., 2001; Lee SK et al., 2004).

In vertebrates, GREs comprise densely clustered TF binding sites which collectively form genomic instructions for the modulation of gene expression and act as switches to turn gene expression on or off (Woolfe et al., 2005; Doh et al., 2007). TFs accumulate to these GREs that can act as enhancers and silencers via specific binding motifs to attract and position the RNA polymerase and additional general TFs to the promoter region for gene activation. This process includes the recruitment of transcriptional co-regulators, mediator complexes, DNA looping and chromatin remodeling for greater DNA access (Blackwood and Kadonaga, 1998; Alberts et al., 2008; Visel et al., 2009). The majority of genes is regulated by complex arrays of enhancers for temporal and spatial expression, so that only when all required TFs are present in a tissue the enhancer becomes active (Alberts et al., 2008; Visel et al., 2009). Furthermore, insulator elements restrict enhancer-promoter interactions to defined chromatin domains (Alberts et al., 2008; Visel et al., 2009). GREs with varying lengths of 50 up to 1500 nucleotides can be located internal to the target gene, as well as up to 1 million base pairs upstream or downstream to the transcription start site, while being able to modulate gene expression independently of their orientation (Khoury and Gruss, 1983; Blackwood and Kadonaga, 1998; Pennacchio et al., 2006; Doh et al., 2007).

Gene coding regions make up a little more than 1% of the human genome, but recent studies revealed that at least 80% of non-coding regions, which were for a long time believed to be evolutionary accumulated ‘junk DNA’, constitute functional sequences...
1. Introduction

(Maher, 2012; Pennisi, 2012). However, in the 1970’s, it has already been predicted that evolutionary changes in anatomy and way of life are more often based on changes in the mechanisms controlling the expression of genes than on changes in protein-coding sequences (King and Wilson, 1975). Biological trait-differences between humans and chimpanzees, for instance, appear to be primarily due to mutations in GREs (Wray and Babbitt, 2008). This hypothesis has been further corroborated, for instance, by the identification of evolutionary rapid changes of only 13 nucleotide substitutions in a limb-specific enhancer that are thought to have been driven by positive selection and that led to drastic gene expression changes with likely consequences for limb evolution (Prabhakar et al., 2008; Wray and Babbitt, 2008). Additionally, changes in GREs are frequently involved in many disease-relevant processes (Visel et al., 2009; Pennisi, 2012). The identification of enhancers is therefore crucial for the understanding of the immense complexity of organisms, yet the responsible information lays somewhere in the ‘infinite deserts’ between the genes proper, which complicates their localization (Nobrega et al., 2003; Maher, 2012).

The ENCODE project, including 30 institutes, 440 scientists and the substantial investment of 185 million USD, aims to catalogue functional DNA sequences and their regulation and function in different cell types for a complex understanding of gene control. After almost 10 years of research with state-of-the-art techniques the participating groups uncovered 70,000 promoter and 400,000 distant enhancer regions (Maher, 2012; Pennisi, 2012). In contrast, hypothesis-driven traditional approaches analyze genomic fragments flanking a gene locus of interest for enhancer activity and narrow down the sequence to the functional enhancer (Uchikawa et al., 2003; Lee SK et al., 2004). In this way, Uchikawa and colleagues identified various specific enhancers driving Sox2 expression in different cell types at different developmental stages, respectively (Uchikawa et al., 2003). Interestingly, the enhancers correspond to conserved non-coding regions between different species (Figure 1.12).

GREs important for accurate expression of vital genes experience selective pressure against change and therefore tend to have a high level of sequence conservation across a wide range of different species (Doh et al., 2007). Accordingly, similarities in sequence between highly divergent organisms imply functional constraint (Woolfe
et al., 2005). Comparison of genomic non-coding sequences reveals a full battery of potential regulatory elements and is a powerful filter to prioritize the search for functional activity sequences in vertebrates (Boffelli et al., 2004; Uchikawa et al., 2004; Uchikawa, 2008; Visel et al., 2008). The server-based genomic alignment tool ‘ECR Browser’ permits fast and automated genome comparison between multiple species which allows pinpointing defined conserved regions as potential GREs within a given genomic locus of interest (Ovcharenko et al., 2004; ecrbrowser.dcode.org).

The identification of enhancers that are capable of driving transcription in specific cells or tissues by combining bioinformatics and experimental tests provides extremely powerful tools for a variety of studies (Timmer et al., 2001; Wray and Babbitt, 2008). Sensory neuron subtype-specific enhancers would facilitate the analysis of the development, function and connectivity of distinct sensory lineages for a better understanding of how the complex somatosensory circuitries develop and function, and how they detect and transduce sensory information (Lallemend and Ernfors, 2012).
1.7. Aim of the study

Despite its decisive importance in somatosensory perception, the relationship between sensory modality, neuronal identity and the stereotypic connectivity of primary sensory neurons remains unclear (Arber, 2012). The available information regarding the mechanisms of molecular specification, precise connectivity and transduction of the many physiologically and morphologically defined sensory neuron subclasses that collectively relay mechanosensory input, is limited due to the lack of available molecular markers (Lumpkin and Caterina, 2007; Marmigère and Ernfors, 2007; Bourane et al., 2009; Luo et al., 2009; Ma, 2009; Lallemend and Ernfors, 2012).

The development of genetic tools for tagging and manipulation of discrete neuronal populations initiated a methodological revolution in virtually all areas of neurobiology (Zhang et al., 2007; Lichtman and Sanes, 2008; Goulding, 2009; O’Conner et al., 2009). In mouse, for instance, gene targeting and bacterial artificial chromosome transgenesis have led to the molecular identification of discrete sensory neuron subtypes and circuitries by mapping of their developmental lineage (Mombaerts, 1996; Zylka et al., 2005; Kim et al., 2008; Bourane et al., 2009; Luo et al., 2009; Chandrashekar et al., 2010). An alternative approach is the selective screen of candidate gene activities putatively linked to given neuronal identities based on data mapped by individual researchers or public gene expression catalogues (Yin et al., 2002; Alvarez-Bolado and Eichele, 2006; Jones et al., 2009). This information can be used to isolate individual gene regulatory elements (GREs) of the candidate genes, to generate transgenic animals in which, hopefully, the desired neuronal population is labeled (Uchikawa et al., 2003; Boffelli et al., 2004; Ovcharenko et al., 2004; Visel et al., 2008). Therefore, the combination of gene expression database screening with GRE identification via genomic comparison between multiple species can potentially reveal sensory subtype-specific enhancers that, in turn, facilitate the analysis of the development, function and connectivity of distinct sensory lineages.

Rodent transgenic studies, however, implicate high costs, resource-intensiveness and time-consuming generations that can be prohibitive to individual researchers for performing neuron subtype-specific GRE screens on a larger basis (Timmer et al.,
1. Introduction

2001). As a complementary model system, the straightforward and rapid in vivo transfection procedures developed in the chick (Gallus gallus domesticus) permit accelerated development of neuron subtype-specific genetic tools and have been proven extremely powerful for uncovering regulatory principles involved in early neural development of both avians and mammals (Novitch et al., 2001; Thaler et al., 2002; Dasen et al., 2005). The chicken is expected to share many features of developmental regulation and shows extensive homology in nervous system organization with mammals, indicating universal conserved gene regulatory mechanisms among vertebrates (Muramatsu et al., 1997; Lee SK et al., 2004; Uchikawa, 2008). Especially circuits involved in somatosensory perception, including morphologically and physiologically identified sensory neurons and their connectivity patterns, are highly conserved between birds and mammals (Necker, 1985; Necker, 1990; Woodbury and Scott, 1991; Duc et al., 1993; Eide and Glover, 1997; Koltzenburg and Lewin; 1997; Guo et al., 2011). Therefore, the in ovo electroporation approach greatly facilitates the study of sensory neuron specification and connectivity in vertebrates.

The initial aim of this work was the identification of genetic markers for distinct sensory subtypes by candidate gene screening and GRE identification via genomic comparison, combined with the establishment of an enhancer activity analysis assay in late-gestation chick embryos. Ultimately, the aim of this work was to evaluate and characterize the potential role of candidate genes in the function and circuit formation of distinct sensory lineages in the chick. The discovery of more restricted genetic markers provides more profound knowledge regarding the sensory system organization and function by linking molecularly identified neuronal subtypes with their preferred sensory modality and their peripheral and central connectivity, which together will allow uncovering of the neural substrates of somatosensation, including touch perception.
2. Materials and methods

2.1. Laboratory consumables and plastic ware

Consumables and plastic ware were purchased from Eppendorf AG, Starlab GmbH and Sarstedt AG. Dissection instruments were obtained from Fine Science Tools GmbH.

2.1.1. Antibodies

<table>
<thead>
<tr>
<th>Antigen</th>
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<tr>
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<td>Goat</td>
<td>1:200</td>
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2. Materials and methods

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<th>Supplier</th>
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<td>A-11122 Life Technologies GmbH</td>
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<tr>
<td>GFP</td>
<td>Sheep</td>
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<tr>
<td>Alexa Fluor 647 conjugate toxin</td>
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**Secondary antibodies**

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</tr>
<tr>
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<td>Donkey</td>
<td>1:2000</td>
<td>Life Technologies GmbH</td>
</tr>
<tr>
<td>Goat</td>
<td>Alexa fluorescent dye</td>
<td>Donkey</td>
<td>1:2000</td>
<td>Life Technologies GmbH</td>
</tr>
<tr>
<td>Sheep</td>
<td>Alexa fluorescent dye</td>
<td>Donkey</td>
<td>1:2000</td>
<td>Life Technologies GmbH</td>
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<tr>
<td>Guinea Pig</td>
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<td>1:500</td>
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**2.1.2. Enzymes**

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<td>Phusion High Fidelity DNA Polymerase</td>
<td>New England Biolabs GmbH</td>
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<td>Platinum Taq DNA Polymerase High Fidelity</td>
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<tr>
<td>Restriction enzymes</td>
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<tr>
<td>Shrimp Alkaline Phosphatase</td>
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<tr>
<td>SP6 RNA Polymerase</td>
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<tr>
<td>T4 DNA Ligase</td>
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<td>T4 Polynucleotide Kinase</td>
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<td>T7 RNA Polymerase</td>
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### 2.1.3. Kits

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<td>QIAprep Spin Miniprep Kit</td>
<td>Qiagen GmbH</td>
</tr>
<tr>
<td>QIAquick Gel Extraction Kit</td>
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<tr>
<td>TOPO TA Cloning Kit</td>
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### 2.1.4. Chemicals and reagents

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<td>Ampicillin</td>
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<td>B-27 supplement</td>
<td>Life Technologies GmbH</td>
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<td>BDNF</td>
<td>R&amp;D Systems Inc</td>
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<td>Benzyl benzoate</td>
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<td>Blocking reagent</td>
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<tr>
<td>Ethanol 99.9%</td>
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### 2. Materials and methods

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<td>Ethyleneglycoltetraacetic acid (EGTA)</td>
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<td>GDNF</td>
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<td>L-15 medium Leibowitz</td>
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<td>Maleic acid</td>
<td>Carl Roth GmbH</td>
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<tr>
<td>Methanol</td>
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<td>NBT/BCIP tablets</td>
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<td>Neurobasal medium</td>
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<td>NGF</td>
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<td>OCT embedding medium</td>
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<td>Poly-L-Lysine</td>
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<tr>
<td>Potassium hydroxide</td>
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2. Materials and methods

Proteinase K AppliChem GmbH
Salmon sperm DNA AppliChem GmbH
Sodium chloride Carl Roth GmbH
Sodium citrate Carl Roth GmbH
Sodium dodecyl sulfate (SDS) Carl Roth GmbH
Sodium hydroxide Carl Roth GmbH
Sucrose Carl Roth GmbH
Tris Carl Roth GmbH
Triton X-100 Carl Roth GmbH
TRIzol reagent Life Technologies GmbH
Trypsin Life Technologies GmbH
Tryptone Carl Roth GmbH
Tween-20 Carl Roth GmbH
Yeast extract AppliChem GmbH

2.1.5. Solutions

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<th>Solution</th>
<th>Contents</th>
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<tr>
<td>Antibody staining solution</td>
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<td>BABB clearing solution</td>
<td>66% Benzyl benzoate; 34% Benzyl alcohol</td>
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<tr>
<td>Culture medium</td>
<td>Neurobasal medium; 1x B-27, 0.5 mM L-Glutamic acid; 25 mM L-Glutamine; 1x Penicillin-Streptomycin</td>
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<tr>
<td>Dent’s solution</td>
<td>80% Methanol; 20% DMSO</td>
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<tr>
<td>Growth factor solution</td>
<td>PBS pH 7.2; 100 µg/ml growth factor</td>
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<tr>
<td>Growth medium</td>
<td>DMEM-F12; 2 µM L-Glutamine; 8 mg/ml Glucose, 2x Penicillin-Streptomycin</td>
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<tr>
<td><em>In situ</em> blocking buffer</td>
<td>MBST pH 7.5; 2% [w/v] Blocking reagent</td>
</tr>
<tr>
<td><em>In situ</em> hybridization buffer</td>
<td>50% Formamide; SSC 5x pH 4.5; 1% SDS; 10 mg/ml Heparin; 10 mg/ml salmon sperm DNA</td>
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<tr>
<td><em>In situ</em> wash buffer I</td>
<td>50% Formamide; SSC 5x pH 4.5</td>
</tr>
<tr>
<td><em>In situ</em> wash buffer II</td>
<td>50% Formamide; SSC 2x pH 4.5</td>
</tr>
<tr>
<td><em>In situ</em> wash buffer III</td>
<td>50% Formamide; SSC 1x pH 4.5</td>
</tr>
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</table>
2. Materials and methods

Laminin solution  D-PBS pH7.2; 1 mg/ml Laminin
LB agar  LB medium; 1% [w/v] Agar-Agar
LB medium  5 g/l Sodium chloride; 5 g/l Yeast extract; 10 g/l Tryptone
Lysis buffer  0.1 M Tris pH 8; 30 mM Sodium chloride; 0.2 mM EDTA;
              0.05% SDS; 0.5 mg/ml Proteinase K
MBST  0.1 M Maleic acid; 0.15 M Sodium chloride; pH 7.5
Mounting solution  PBS pH 7.2; 50% Glycerol
NTMT  100 mM Sodium chloride; 10 mM Tris-Hydrochloric acid;
       50 mM Magnesium chloride; pH 9.5
PBST  PBS pH 7.2; 0.1% Tween-20
PFA solution  PBS pH 7.2; 4% [w/v] Paraformaldehyde
PFA culture solution  PBS pH 7.2; 2% [w/v] Paraformaldehyde; 15% [w/v] Sucrose
SSC 20x solution  3 M Sodium chloride; 300 mM Sodium citrate; pH 4.5
Sucrose solution  PBS pH 7.2; 30% [w/v] Sucrose
TAE buffer  40 mM Tris; 1 mM EDTA adjust pH 7.5 with glacial acetic acid

2.1.6. Vectors

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2.1.7. Software

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<td>Technelysium Pty Ltd</td>
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<td>ECR Browser</td>
<td>ecrbrowser.dcode.org/</td>
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<tr>
<td>ImageJ</td>
<td>rsbweb.nih.gov/ij/</td>
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<tr>
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<tr>
<td>pDraw</td>
<td>Acaclone Inc</td>
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2.2. Molecular biology

The methods described were performed according to ‘Molecular cloning: a laboratory manual’ (Sambrook and Russell, 2001) unless otherwise stated.

2.2.1. Polymerase chain reaction

The amplification of DNA fragments was performed with the Platinum Taq DNA Polymerase High Fidelity, the Phusion High Fidelity DNA Polymerase or the Dream Taq DNA Polymerase using the Mastercycler epGradient S (Eppendorf AG). The reaction mixture and the PCR program were used according to the manufacturer’s protocols. Amplified DNA fragments were analyzed via agarose gel electrophoresis.

2.2.2. DNA restriction digest

The digestion of DNA was performed by the use of appropriate restriction enzymes (Thermo Fisher Scientific GmbH). The DNA was incubated with the enzymes and the reaction buffers at optimal reaction temperatures recommended by the manufacturer’s reference for at least 3 h. Double digestions were carried out with reaction buffers enabling maximal enzyme activity for both endonucleases. Alternatively the digested DNA was purified with the QIAquick Gel Extraction Kit (Qiagen GmbH) before the second restriction digest.
2.2.3. Alkaline phosphatase treatment
In order to prevent religation, the vector DNA was dephosphorylated by adding Shrimp Alkaline Phosphatase (Thermo Fisher Scientific GmbH) to the completed restriction digest of the vector DNA and incubation for 30 min at 37°C.

2.2.4. Agarose gel electrophoresis
The separation of DNA fragments was performed by agarose gel electrophoresis. Therefore 1% [w/v] agarose was dissolved in TAE buffer by heating, followed by adding 0.05 µl/ml of ethidium bromide solution and transferring in a gel tray. After solidification, the gel was inserted into a gel chamber covered with TAE buffer, the DNA samples were mixed with DNA loading dye and loaded into the gel pockets. The electrical current was applied until a conclusive segregation of the DNA fragments was achieved.

2.2.5. DNA purification from agarose gels
After the agarose gel electrophoresis, DNA bands were visualized under UV light illumination and the appropriate DNA fragments were extracted from the agarose gel. They were purified by using the QIAquick Gel Extraction Kit (Qiagen GmbH) according to the manufacturer's instructions.

2.2.6. DNA ligation
The ligation mixture of digested vector and insert DNA fragments was prepared with a molar concentration ratio of 1:4. Therefore, the DNA concentrations were determined and incubated with T4 DNA Ligase and the appertaining buffer (Thermo Fisher Scientific GmbH) at 22°C for at least 6 h or with the DNA Ligation Kit (Takara Bio Inc) at 16°C for at least 4 h. For PCR products with low concentrations, the TOPO TA Cloning Kit (Life Technologies GmbH) was used according to the manufacturer's instructions.
2. Materials and methods

2.2.7. Plasmid preparation
Single bacteria colonies were picked from agar plates and incubated in 5 ml LB medium containing ampicillin overnight at 37°C. A plasmid minipreparation was performed using the QIAprep Spin Miniprep Kit (Qiagen GmbH) due to the manufacturer’s instructions. Positive clones were verified via restriction digest and sequencing. For a plasmid maxipreparation, the *E.coli* culture was grown from 100 µl of a glycerol stock in 200 ml LB medium with ampicillin overnight at 37°C and the plasmid DNA was extracted with the QIAGEN Plasmid Maxi Kit according to the manufacturer’s manual.

2.2.8. DNA concentration measurements
The DNA and RNA concentration was determined using the Nanodrop 1000 UV-Vis Spectrophotometer (Thermo Fisher Scientific GmbH) based on UV light absorption of the nucleic acids at definitive wave lengths.

2.2.9. Sequencing
DNA sequencing was carried out by the Qiagen genomic services (Qiagen GmbH). DNA samples were submitted in a concentration of 500 ng per reaction. Only plasmids with strictly accurate sequences were used for further experiments.

2.2.10. Preparation of chemically competent *E.coli*
Chemically competent *E.coli* cells of the One Shot TOP10 strain (Life Technologies GmbH) were used for all molecular cloning experiments. The preparation was performed according to Inoue *et al.* (Inoue *et al.*, 1990) and stored at -80°C until use.

2.2.11. Transformation
Chemically competent *E.coli* were transformed by adding of 100 µl of the soluble cell suspension to the ligation mixture. After 30 min of incubation on ice, the cells were heat shocked at 42°C for 1 min and chilled on ice afterwards. *E.coli* were grown in 1 ml
2. Materials and methods

of antibiotic-free LB medium at 37°C for one hour on a shaker and plated on an agar plate containing ampicillin. The plates were incubated overnight at 37°C.

2.2.12. Preservation and recovery of \textit{E.coli}

The preservation of the \textit{E.coli} clones was performed by mixing 500 µl of soluble \textit{E.coli} culture with 500 µl glycerol in a reaction tube and storing of the mixture at -80°C. For regrowing, 100 µl of the frozen \textit{E.coli} stocks were incubated in 5 ml LB medium with ampicillin for a plasmid minipreparation or in 150 ml LB medium with ampicillin for a plasmid maxipreparation overnight at 37°C.

2.2.13. DNA extraction

Genomic DNA of mouse and chicken for enhancer amplification was extracted from mouse tail tip biopsies and whole chicken embryo of embryonic day 4 (E4) by incubation in lysis buffer overnight at 56°C. The lysate was cleared by centrifugation and the supernatant was mixed with 500 µl isopropanol. After centrifugation, the DNA pellet was washed with 500 µl 70% ethanol, air dried and eluted in water.

2.2.14. RNA extraction

Total RNA from mouse spinal cords and dorsal root ganglia (DRGs) from embryonic day 18.5 (E18.5) and chicken DRGs from embryonic day 12 (E12) was obtained via TRizol isolation. The tissue was homogenized in 1 ml TRizol reagent per 100 mg of tissue and incubated for 5 min. The samples were incubated with 0.2 ml chloroform per 1 ml TRizol for 3 min and centrifuged for 15 min at 4°C. The aqueous phase containing the RNA was removed, transferred into a new reaction tube and incubated with 0.5 ml isopropanol per 1 ml of TRizol and 3 µl of polyacrylamide for 10 min. After centrifugation for 15 min at 4°C, the supernatant was removed and the RNA pellet was washed with 1 ml 75% DEPC-ethanol. The samples were centrifuged, air dried and eluted in DEPC-H₂O.
2.2.15. cDNA synthesis
The cDNA for the gene amplification and the in situ probe generation was synthesized using the iScript cDNA synthesis Kit (Bio-Rad Laboratories GmbH) or the PrimeScript First Strand cDNA Synthesis Kit (Takara Bio Inc) according to the manufacturer’s instructions.

2.3. Plasmid construction

The plasmid pSTEVE was designed for reporter gene-based screening of cis-regulatory activities, combined with Tol2 transposon-based transgenesis facilitating genomic integration upon in ovo electroporation (Kawakami and Shima, 1999; Sato et al., 2007; Takahashi et al., 2008). The plasmid pSTEVE-eGFP was constructed by insertion of the Tol2 sites from pT2KXIGΔin vector in the pUC18 vector, followed by insertion of a minimal TATA box promoter and eGFP from pGATA2V2X2 between the Tol2 sites. The eGFP reporter gene was replaced by the mGFP gene from pCAG-mGFP to generate pSTEVE-mGFP. The putative enhancers were subcloned upstream of the minimal promoter which do not drive a reporter gene expression without an additional enhancer. Isl1Crest3 was amplified from mouse genomic DNA with following primers: 5’-TAAAAGAGCAAACTACAGC-3’ and 5’-AATAGTGTTTCTTGGC-3’. AvilLucy1 was amplified from mouse genomic DNA with following primers: 5’-GGGAATTGGAGGCTCCTG-3’ and 5’-ACCCTCTGTCCTCCGAAGT-3’. Ntng1Mech1 was amplified from chick genomic DNA with following primers: 5’-GATCCTTAATGTGCTTTGGT-3’ and 5’-AGTCATAATGATGGTCGT-3’. TrkAECR1 was amplified from mouse genomic DNA with following primers: 5’-GCTCGCCTTCCTCTCATTGTC-3’ and 5’-CTCAGGATGCTTAGGTTG-3’. Runx3ECR1 was amplified from mouse genomic DNA with following primers: 5’-TAATAAATGGCCCGAGAA-3’ and 5’-CTCAGGATGCTTAGGTTG-3’. Runx3ECR2 was amplified from mouse genomic DNA with following primers: 5’-TTGTGTACCAGACAGGAGCA-3’ and 5’-CCACTCTCCTGCTGCTCTAA-3’. Runx3ECR3 was amplified from mouse genomic DNA with following primers: 5’-CAGTCGGCTGGACCTGAA-3’ and
2. Materials and methods

5’-TTGCTGCCCTTGACCTTT-3’. The CMV early enhancer / chicken β-actin (CAG) promoter was cut out from pCAG-mGFP via the XbaI and SpeI restriction sites. The created plasmids were verified via sequencing. pSTEVE-eGFP was used to visualize the cell somata of electroporated (dorsal root ganglion) DRG neurons, and pSTEVE-mGFP to trace sensory axons. The transposase in the pCAGGS-T2TP was co-electroporated for a stable genetic integration in the transfected cells (Figure 2.1).

For overexpression of Ntng1 selectively in sensory neurons and the ligand NGL1 in the entire spinal cord, the Cre/loxP system was used to combine a high expression level of the gene with maintaining the specificity in sensory neurons or the spinal cord (Livet et al., 2007). Therefore the pCAG-loxP-ssFlag-IRES-mGFP/tdTomato-Tol2 vector was designed consisting of six essential parts: the strong ubiquitous CAG promoter (1), the loxP-STOP-loxP cassette (2) for Cre recombination, followed by a signal sequence (3), which is necessary for the secretion pathway and a cellular membrane insertion, an Flag epitope tag (4) upstream of the gene insertion sites for protein detection and reporter genes IRES-mGFP or IRES-tdTomato (5). The expression cassette was flanked by Tol2 sites (6) for transposase-mediated stable genomic integration.

The Ntng1 coding sequence without its signal sequence was amplified from cDNA of mouse embryonic day 18.5 (E18.5) DRGs and inserted downstream of the Flag tag peptide sequence into the pCAG-loxP-ssFlag-IRES-mGFP-Tol2 construct. The following primers were used: 5’-GTGATGCAGCCCTACCTTTTCG-3’ and 5’-CTAGAACACCAGGGGACCAGCA-3’. The NGL1 coding sequence without its signal sequence was amplified from cDNA of mouse embryonic day 18.5 (E18.5)
spinal cord and inserted downstream of the Flag tag peptide sequence into the \( pCAG-loxP-ssFlag-IREs-tdTomato-Tol2 \) construct. The following primers were used: 5'-CAAACCTGCCCTTCAGTGTGCT-3' and 5'-TTATATCTGAGTCTCTTGTACA-3'. The sensory-specific \( \text{pIsl1Crest3-Cre} \) driver plasmid was created by insertion of a minimal TATA box promoter and eGFP from \( p\text{GATA2V2X2} \) in the \( p\text{UC18} \) vector, following the replacement of the eGFP gene by the Cre gene from the \( p\text{PGK-Cre-bpA} \) vector and insertion of the \( \text{Isl1Crest3} \) enhancer upstream of the minimal promoter. For the expression of NGL1 in the spinal cord, the constitutive \( p\text{PGK-Cre-bpA} \) vector was used. The final plasmids were verified via sequencing. The transposase in the \( p\text{CAGGS-T2TP} \) was co-electroporated for a stable genetic integration in the transfected cells (Figure 2.2).

### Figure 2.2: Plasmid construction for transgene expression analysis

For overexpression of NGL1 in the spinal cord, the gene was cloned into the expression vector and co-electroporated with the general Cre driver (\( p\text{PGK-Cre} \)) and the transposase for stable genomic integration. Overexpression of Ntng1 was achieved by subcloning of the gene coding sequence into the expression vector and co-electroporation with the sensory-specific \( \text{pIsl1Crest3-Cre} \) driver and the transposase for stable genomic integration.

### 2.4. In ovo electroporation

Fertilized chick eggs (Horstmann Geflügelzucht GmbH) were incubated at 38°C and 80% humidity. The in ovo electroporation of DNA constructs was performed between Hamburger-Hamilton (HH) stages 11 and 13 (embryonic day 2 - 2.2) using the ECM 830 electroporation system (BTX Instrument Division, Harvard Apparatus Inc) with
2. Materials and methods

Following settings: 5 pulses of 25 mV for 50 ms in the LV 99 ms/500 V modus. All plasmids were injected into the central canal of the neural tube at final concentrations of 1.5 µg/µl in TE in a 4:1 mixture with the pCAGGS-T2TP plasmid. For overexpression the molar ratio for expression vector, Cre driver vector and transposase vector was 2:1:1. in the chick (embryonic day 2 - 2.2). Electroporated embryos were incubated until appropriate embryonic stages. All animal work has been conducted according to national and international guidelines.

2.5. Immunohistochemistry

Chicken spinal cords with attached dorsal root ganglia (DRGs), as well as hindlimbs and digits were dissected at appropriate developmental stages, fixed at 4°C in PFA solution, washed in PBS, immersed in sucrose solution and embedded into OCT embedding medium according to the incubation timeline (Table 2.1). 60 µm transversal cryosections were cut in a CM1900 cryostat (Leica Microsystems GmbH) at −20°C, placed on a super frost glass slide and stored at -20°C. For primary antibody detection the slides were washed three times for 30 min with PBS and incubated overnight at 4°C in staining solution. Slides were washed three times for 30 min with PBS and stained with secondary antibody solution for 1 h at room temperature. The slides were washed with PBS, mounted with mounting solution and cover slipped.

Table 2.1: Incubation timeline for chicken embryos

<table>
<thead>
<tr>
<th></th>
<th>E4</th>
<th>E5</th>
<th>E6</th>
<th>E7</th>
<th>E9</th>
<th>E12</th>
<th>E15</th>
<th>E19</th>
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<td>1 h</td>
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<td>4 h</td>
<td>6 h</td>
<td>10 h</td>
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<td>PBS</td>
<td>5 h</td>
<td>6 h</td>
<td>7 h</td>
<td>8 h</td>
<td>10 h</td>
<td>12 h</td>
<td>15 h</td>
<td>20 h</td>
</tr>
<tr>
<td>Sucrose</td>
<td>3 h</td>
<td>4 h</td>
<td>5 h</td>
<td>6 h</td>
<td>7 h</td>
<td>8 h</td>
<td>10 h</td>
<td>15 h</td>
</tr>
</tbody>
</table>

Whole mounts of embryonic day 5 (E5), whole hindlimbs of embryonic day 9 (E9) and whole digits of embryonic day 15 (E15) were fixed in PFA solution for 5 h, washed in PBS overnight, bleached with Dent’s solution for 5 h, rehydrated in PBST overnight.
and incubated in primary antibody solution for 4 days. The limbs were washed in PBST overnight, incubated in secondary antibody solution for 4 days, washed in PBST overnight, immersed in a graded series of methanol solution up to 100% and cleared with BABB solution. All steps were carried out at 4°C.

2.6. *In situ* hybridization

The *in situ* hybridization probe for the chicken Ntng1 coding sequence was generated by PCR amplification of a 654 bp fragment from cDNA of chicken embryonic day 9 (E9) dorsal root ganglia (DRGs) by the use of following primers: 5’-ATGATGTATTTGTCGAGATTTCT-3’ and 5’-ACTGTTTGTCATATATCCAG-3’. The probe was T/A subcloned into the pCRII-TOPO vector and verified via sequencing. The antisense RNA probe was synthesized by the T7 or Sp6 RNA polymerase using the DIG RNA labeling Kit according to the manufacturer’s specifications. The quality and quantity was verified.

Chicken spinal cords with attached DRGs at different developmental stages were dissected, collected in D-PBS, fixed in PFA solution for 48 h, washed in D-PBS for 24 h and dehydrated in 30% sucrose solution. All steps were carried out at 4°C. The samples were embedded into OCT embedding medium, cryosectioned at 12 µm in a CM1900 cryostat (Leica Microsystems GmbH) at −20°C and placed on a super frost glass slide.

For the *in situ* hybridization, the slides were washed with D-PBS for 3 min, dehydrated in a graded series of methanol in D-PBS and stored in 100% methanol at -20°C overnight. The samples were rehydrated through methanol in D-PBS series for 5 min, bleached in 6% hydrogen peroxide in D-PBS and washed three times in D-PBS for 5 min. The sections were treated with 10 µg/ml Proteinase K in D-PBS for 10 min, washed with D-PBS and postfixed with PFA solution for 5 min. They were washed with D-PBS for 5 min and 0.85% NaCl in D-PBS, followed by dehydration through graded series of ethanol solutions for 10 min and air drying. The samples were transferred in a
2. Materials and methods

Humidified chamber containing *in situ* wash buffer I and prehybridized with *in situ* hybridization buffer for one hour at 68°C. The slides were treated with *in situ* hybridization buffer containing 500 ng/ml of the DIG-labeled antisense RNA probe, covered with parafilm and incubated overnight at 68°C. After hybridization, the parafilm was removed and the samples were immersed in *in situ* wash buffer I-III at 65°C for 30 min. The slides were washed two times with MBST at 70°C for 30 min and three times with MBST at room temperature for 5 min, before they were treated with *in situ* blocking buffer at room temperature for one hour. The samples were transferred in a humidified chamber containing water and incubated with 1:2500 diluted anti-digoxigenin alkaline phosphatase conjugated antibody in *in situ* blocking buffer overnight at 4°C. The slides were washed three times with MBST for 5 min and treated with NTMT for 10 min. The *in situ* signal was developed by incubation in NBT/BCIP dissolved in water at room temperature, followed by washing with D-PBS and mounting with mounting solution.

2.7. Cell culture

For the DRG explant culture assay, transfected embryonic day 6 (E6) chicken dorsal root ganglia (DRGs) from lumbar spinal segments were dissected, collected in culture medium on ice and transferred in a four-well plate on a glass coverslip precoated with 1 mg/ml Poly-D-Lysine and 100 µg/ml Laminin (Wang and Marquardt, 2012). BDNF, GDNF, NGF or NT-3 growth factor medium was added to the culture medium and the plate was transferred into the incubator overnight.

For immunohistochemistry, the explant samples were fixed by adding the equal amount of PFA culture solution to the culture medium for 30 min. After removing the top phase of the solution, another volume of the fixative was added for 1 h. The samples were washed three times for 10 min with PBST and incubated in primary antibody staining solution overnight at room temperature. After three washing steps with PBST, the secondary antibody solution was applied for 1 h at room temperature.
The samples were washed with PBST and mounted by inverting the coverslip with the explants on the mounting solution coated microscope slide.

For electrophysiological measurements, transfected embryonic day 9 (E9) chicken DRGs from all spinal segments were dissected, collected in Ca\(^{2+}\) and Mg\(^{2+}\)-free PBS and treated with 0.05% trypsin for 12 min at 37°C. Digested DRGs were washed twice with growth medium, triturated and plated in a droplet of growth medium on a glass coverslip precoated with 20 µg/cm\(^2\) Poly-L-Lysine and 4 µg/cm\(^2\) Laminin. Neurons were allowed to adhere to the coverslips for 3-4 hours at 37°C in a humidified 5% incubator before being used for patch-clamp experiments.

The DRG culture for electrophysiological measurements was performed in collaboration with Stefan G. Lechner and Gary R. Lewin at the ‘Max-Delbrück-Center for Molecular Medicine’ in Berlin.

### 2.8. Electrophysiology

Whole cell patch clamp recordings were made at room temperature from cultures prepared as described above. Patch pipettes were pulled from borosilicate glass capillaries, filled with a solution consisting of 110 mM KCl, 10 mM NaCl, 1 mM MgCl\(_2\), 1 mM EGTA and 10 mM HEPES, adjusted to pH 7.3 with KOH and had tip resistances of 2-4 MΩ. The bathing solution contained 140 mM NaCl, 4 mM KCl, 2 mM CaCl\(_2\), 1 mM MgCl\(_2\), 4 mM Glucose, 10 mM HEPES, adjusted to pH 7.4 with NaOH. All recordings were made using an EPC 10 amplifier (HEKA Elektronik Dr. Schulze GmbH) in combination with the Patchmaster and Fitmaster software (HEKA Elektronik Dr. Schulze GmbH). Pipette and membrane capacitance were compensated using the auto function of Patchmaster and series resistance was compensated by 70% to minimize voltage errors.

Action potentials were evoked by repetitive 80 ms current injections increasing from 40 pA to 800 pA in increments of 40 pA. The first action potential evoked with this
2. Materials and methods

A pulse protocol was used for analysis. Mechanically activated currents were recorded as previously described (Hu and Lewin, 2006; Lechner et al., 2009). Briefly, neurons were clamped to -60 mV, stimulated mechanically with a fire-polished glass pipette (tip diameter 2-3 µm) that was driven by a piezo based micromanipulator called nanomotor (Kleindiek Nanotechnik GmbH) and the evoked whole cell currents were recorded with a sampling frequency of 200 kHz. The stimulation probe was positioned at an angle of 45° to the surface of the dish and moved with a velocity of 3.5 µm/ms. Currents were fitted with single exponential functions and classified as RA-, IA- and SA-type currents according to their inactivation time constant (Hu and Lewin, 2006). All electrophysiological experiments were carried out on acutely dissociated dorsal root ganglion (DRG) neurons between 3 and 8 hours after plating.

The electrophysiological measurements were performed in collaboration with Stefan G. Lechner and Gary R. Lewin at the ‘Max-Delbrück-Center for Molecular Medicine’ in Berlin.

2.9. Imaging

All images, including those of transverse sections and whole mounts, were collected using a Leica TCS/MP confocal/two-photon microscope (Leica Microsystems GmbH) with 488 nm, 543 nm and 633 nm laser lines, in situ hybridization images were taken in the bright field. All images of a stack of ≥18 z-sections subsequently collapsed to a 2D rendering. The pictures were processed using Adobe Photoshop CS5.

2.10. Quantifications and measurements

For the quantification of ipsilaterally and contralaterally migrating Isl1Crest3+, AvilLucy1+ and CAG+ sensory neurons, transverse sections of lumbar dorsal root ganglia (DRGs) of at least 8 embryos were selected. GFP+ / Isl1+ DRG neurons were counted on two
or more non-adjacent sections per embryo and presented as the mean ± standard error of the mean (SEM). Statistical comparisons of cell counts were established using a two-tailed Student's t test with two samples and unequal variance.

For the comparison of Ntng1^{Mech1+} and Isl1^{Crest3+} DRG neurons, GFP^+ cells on transverse lumbar sections of at least 8 embryos were scored for colocalization with TrkA, TrkB, TrkC, Ret or NF200 and analyzed as the mean ± standard error of the mean (SEM). Statistical comparisons of colocalization counts were established using a two-tailed Student's t test with two samples and unequal variance. Additionally the cell size area was measured of GFP^+, TrkA^+, TrkB^+ or TrkC^+ neurons by cell border determination of neurons with clear morphology and visible nucleus.

Projection density maps were generated from 18 sections from the lumbar dorsal horn segments out of ≥ 8 embryos. A stack with average intensity was performed that was rendered as a numerical grid based on pixel intensity values and converted into a normalized heat map. The statistical analysis of the laminae I-II versus laminae III-V axon density was performed by signal intensity comparison of the laminae based on mean pixel intensity values from 18 sections according to the projection density screen and presented as the mean ± SEM by using a two-tailed Student's t test with two samples and unequal variance.
3. Results

3.1. Stable genetic tagging of somatosensory neurons in chick

The assembly of circuitries for somatosensory perception relies on a complex interplay of neuronal diversification, specification, phenotypic modulation and precise connectivity. However, little is known about the molecular mechanisms controlling the establishment of well-defined somatosensory circuitries, which is to a large part due to the limitation of sensory neuron subtype-specific molecular markers (Arber, 2012; Lallemend and Ernfors, 2012). In order to identify novel markers for distinct sensory subtypes, published literature and public gene expression catalogues were selectively screened for candidate genes expressed by subsets of dorsal root ganglion (DRG) neurons. Through this information, potential gene regulatory elements of the corresponding candidate genes, capable of driving transcription in specific cell types, were defined by an \textit{in silico} screen for evolutionary conserved non-coding regions (ECRs) via browser-based genome alignment between multiple species (Ovcharenko \textit{et al.}, 2004; \texttt{ecrbrowser.dcode.org}). The chicken model system shares developmental and structural similarities with mammals, including highly conserved somatosensory system organization and overall connectivity, and allows a more rapid genetic manipulation and analysis than rodent transgenic studies (Necker, 1990; Eide and Glover, 1997; Koltzenburg and Lewin, 1997; Timmer \textit{et al.}, 2001; Guo \textit{et al.}, 2011). The \textit{in ovo} electroporation technique has been widely used for enhancer activity analysis \textit{in vivo} and has proven to be a powerful tool for uncovering regulatory principles involved in early neural development of both avians and mammals (Novitch \textit{et al.}, 2001; Thaler \textit{et al.}, 2002; Dasen \textit{et al.}, 2005; Uchikawa, 2008). Therefore, a straightforward strategy was explored allowing effective, selective and stable genetic tagging of somatosensory neurons and their connectivity patterns in chick embryos (Figure 3.1 a).

For a precise analysis of enhancer activity, especially in late-gestation mature or nearly mature sensory neurons, a stable somatic transfection vector, termed \textit{STEVE} (\textit{STEVE: stable expression vector}), was designed. \textit{STEVE} consists of three essential parts: a basal promoter, a reporter gene and flanking transposable elements. The
identified ECRs corresponding to putative enhancers were amplified from genomic DNA and subcloned upstream of the minimal promoter. The minimal promoter itself should not drive significant own activity in detectable expression levels, but was expected to faithfully reflect the activity of the enhancers contained within inserted ECRs. In addition, the sensitivity of the reporter gene should allow readily accessible detection of low-level expression driven by putative enhancers. By testing different basal promoters and reporter genes for their suitability for \textit{in ovo} enhancer studies, a minimal \textit{TATA box} basal promoter cassette, coupled to reporter genes encoding green fluorescent protein (GFP), proved to be the most appropriate choice to perform a systematic \textit{in ovo} screen for enhancer activities. For different experimental purposes, either cytoplasm-localized \textit{enhanced GFP (eGFP)} or membrane-tethered and axon-localized \textit{myristoylated GFP (mGFP)} were inserted as reporter genes. The technique of \textit{in ovo} transfection of the neural tube, including neural crest cells (NCCs), which give rise to DRG neurons, provides an accessible route for introducing transgenes into the sensory neuron lineage (Muramatsu \textit{et al.}, 1997; Krispin \textit{et al.}, 2010). However, the transient nature of these ‘pseudogenetic’ manipulations so far precluded their application to more mature aspects of sensory nervous system connectivity and function. Since the expression cassettes are not integrated into the host’s genome, their copies become diluted and ultimately disappear as NCCs undergo massive proliferation (Yokota \textit{et al.}, 2011). To overcome these limitations, an effective, stable genetic tagging of transfected cells was achieved by transposon-based transgenesis based on the medaka fish hAT family \textit{Tol2} transposon (Kawakami and Shima, 1999; Sato \textit{et al.}, 2007; Takahashi \textit{et al.}, 2008). The \textit{Tol2} transposase (\textit{T2TP}) recognizes a pair of core transposon sites (\textit{t2}) flanking the ‘enhancer-promoter-reporter gene’ expression cassette, resulting in the recombination of the transgene sequence from the vector and its subsequent random integration in the genome. To achieve this, the transposase driven by the strong and ubiquitously active \textit{CMV early enhancer / chicken β-actin (CAG) promoter} (\textit{pCAGGS-T2TP}) was therefore co-electroporated with \textit{STEVE}.

The efficiency of the designed strategy and the optimal electroporation conditions were first determined by examining the expression of a reporter gene driven by the CAG promoter (\textit{pCAG-STEVE-eGFP}). The vast majority of cells on the transfected
3. Results

(ipsilateral) side of the spinal cord expressed high levels of eGFP at E12 (Figure 3.1 b-c), while no activity was detected in embryos transfected with enhancer-less constructs (Figure 3.1 d-e). The electroporation showed no influence on embryonic development based on overall morphology, as well as neural marker gene expression, including Isl1 protein expression, between the ipsilateral and contralateral side. The in ovo electroporation was performed between Hamburger-Hamilton (HH) stages 11 and 13 in the chick (embryonic day 2 - 2.2) before NCC emigration for a high transfection efficiency of somatosensory neuron progenitors (Hamburger and Hamilton, 1951; Krispin et al., 2010).

Figure 3.1: Selective and stable genetic tagging of spinal and somatosensory neurons in chick

(a) Strategy: Subcloning of in silico identified evolutionary conserved non-coding regions (ECRs) into transposon-based pSTEVE-eGFP or mGFP. Co-electroporation with transposase (T2TP) into the neural tube of a chick embryo HH stage 11-13 (E2 - 2.2) for stable transfection of neural crest cells (NCCs) before their emigration and formation of the dorsal root ganglia (DRGs). Analysis of putative enhancer activities in late-gestation embryos.

(b-c) The constitutive CAG promoter (pCAG-STEVE-eGFP) drives eGFP expression in virtually all cells on the transfected (ipsilateral) side of the chick spinal cord in E12 transverse sections. Isl1 protein expression is restricted to motor neurons and dI3 interneurons. Dotted lines outline spinal cord gray matter. Scale bar: 100 µm.

(d-e) No expression is detected in embryos transfected with enhancer-less constructs (pSTEVE-eGFP) in transverse sections at E12. Scale bar: 100 µm.
Based on literature search and gene expression databases, a set of candidate genes with general and subtype-restricted expression in somatosensory neurons were selected for the in silico-to-in vivo screen and the identified ECRs were analyzed for their spatial and temporal expression pattern in the sensory lineage at both early and late stages of gestation (Table 3.1). The neuron type composition of NCC progeny tagged by stable reporter protein expression was investigated at different developmental stages spanning embryonic day (E) 5 to E15 for all examined ECRs.

<table>
<thead>
<tr>
<th>sensory lineage</th>
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<th>ECR</th>
<th>genomic focus</th>
<th>ECR activity</th>
<th>ECR specificity</th>
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<td>Isl1</td>
<td>CREST3</td>
<td>mm chr13: 116,633-179-116,634,173 995 bps</td>
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<td>yes</td>
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<tr>
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<td>Lucy1</td>
<td>mm chr10: 127,000-229-127,000,828 600 bps</td>
<td>all sensory neurons</td>
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Candidate genes with general and subtype-restricted expression in somatosensory neurons were screened for evolutionary conserved non-coding regions (ECRs). The identified ECRs were analyzed via in ovo electroporation for their spatial and temporal expression activity in late-gestation sensory neurons and for their specificity for the endogenous gene expression of the corresponding genes.

As a general somatosensory marker and a proof of principle experiment, the Isl1 gene was chosen, which at spinal levels is expressed at earliest stage of neural differentiation by all subtypes of somatosensory neurons, somatic and preganglionic motor neurons, as well as by dorsal (dl3) interneurons (Ericson et al., 1992). The transcription factor Isl1 is highly conserved during evolution and plays a central role in the transition from sensory neurogenesis to subtype specification, whereby gene expression is mediated by different subtype-specific enhancers (Tsuchida et al., 1994;
Thaler et al., 2004; Uemura et al., 2005; Sun et al., 2011). Herein, an ECR at around 323 kb upstream of the mouse Isl1 gene transcription start site coincides with a genomic fragment previously been shown to drive expression in sensory neurons in zebrafish (Uemura et al., 2005). This mouse genomic ECR, Isl1\textsuperscript{Crest3}, was found to drive specific and robust reporter gene expression in sensory DRG neurons of transfected chick embryos (see chapter 3.2).

In order to establish a late-gestation and adult pan-sensory marker, the Advillin (Avil) gene locus was screened for a putative enhancer. The actin binding protein Avil, which is highly conserved between mammals but not yet identified in the chick genome, is specifically expressed in sensory neurons and implicated in regenerative neurite outgrowth (Hasegawa et al., 2007; Lau et al., 2011; Zurborg et al., 2011). The identified ECR Avil\textsuperscript{Lucy1} locates just upstream to the gene coding sequence. Upon isolation, subcloning into STEVE and in ovo electroporation, Avil\textsuperscript{Lucy1} was able to drive a robust expression highly restricted to sensory DRG neurons, which was retained at least until hatching (see chapter 3.3).

The growth factor receptor Tropomyosin receptor kinase A (TrkA) is confined to be expressed in most pain-transmitting neurons and plays an important role in shaping the nociceptive phenotype (Martin-Zanca et al., 1990; Smeyne et al., 1994; Ma et al., 2003). A core enhancer located upstream adjacent to the gene transcription start site was shown to drive somatosensory-specific expression in mouse (Ma et al., 2000). This TrkA\textsuperscript{ECR1}, which was not conserved in the chick genome, was tested for enhancer activity in the chick embryo. However, no reporter gene expression could be observed in DRG neurons at different developmental stages (data not shown; see chapter 4.2 for discussion).

Proprioceptive neurons highly express the Runt-related transcription factor 3 (Runx3) as a key factor in their specification and functional circuit formation (Inoue et al., 2002; Levanon et al., 2002; Chen AI et al., 2006). Three ECR sequences with a conservation of around 80% between mouse and chicken were selected for enhancer analysis spanning a region from 95 kb to 31 kb upstream of the gene transcription start site. Both Runx3\textsuperscript{ECR1} and Runx3\textsuperscript{ECR2} were driving expression in sensory neurons which
appeared more wide-spread and not restricted to the proprioceptive subtype, whereas Runx3\textsuperscript{\text{ECR3}} did not show a detectable expression in transfected cells at different developmental stages (data not shown). Therefore, the identified ECRs did not reflect the endogenous Runx3 gene expression and were not characterized further (see chapter 4.2 for discussion).

NetrinG1 (Ntng1) appears to be expressed by a subset of large-diameter sensory neurons in postnatal mouse DRGs and encodes a Glycosyl-phosphatidyl-inositol (GPI)-linked interaction partner of the transmembrane NetrinG1 ligand (NGL1), which is implicated in laminar neurite targeting and synaptogenesis (Yin et al., 2002; www.brain-map.org). Although the role of Ntng1 in somatosensory neurons remains unknown, it might be involved in sensory connectivity (Nakashiba et al., 2002; Nishimura-Akiyoshi et al., 2007; Woo et al., 2009). An identified ECR located 2.6 kb upstream of the gene coding sequence showed a specific expression in a subclass of large-diameter mechanoreceptive neurons (see chapter 3.5).

Alltogether, the designed strategy, including the \textit{in silico}-to-\textit{in vivo} ECR screen and the stable expression system STEVE, thus facilitates the detection of cis-regulatory activities and their subsequent use for the stable genetic tagging of somatosensory neurons in chick.
3. Results

3.2. Analysis of the sensory neuron-specific enhancer \textit{IsI1}^{Crest3}

During early vertebrate embryogenesis, the transcription factor Islet1 is expressed in a defined population of somatosensory neurons, somatic and preganglionic motor neurons, as well as dorsal (d13) interneurons (Ericson \textit{et al.}, 1992). Thereby, the evolutionary highly conserved factor has been shown to be critical in neuronal specification of both motor and sensory neurons (Tsuchida \textit{et al.}, 1994; Thaler \textit{et al.}, 2004; Sun \textit{et al.}, 2011). The endogenous gene expression is mediated by distinct neuron type-specific enhancers (Uemura \textit{et al.}, 2005). A through \textit{in silico} screen identified ECR, \textit{IsI1}^{Crest3}, at around 323 kb upstream of the mouse \textit{IsI1} coding sequence coincided with a genomic fragment previously been shown to predominantly drive expression in sensory neurons in zebrafish (Figure 3.2 a) (Uemura \textit{et al.}, 2005). This mouse ECR with a length of 995 bps revealed 91% and 85% homologies with the human and chicken sequences, respectively, and included 94 putative conserved transcription factor binding sites (TFBS) (Figure 3.2 b).

Based on this information, \textit{IsI1}^{Crest3} was chosen as a putative early stage pan-sensory marker and analyzed for the spatial and temporal enhancer activity in the chick somatosensory lineage. When coupled to \textit{STEVE \textit{IsI1}^{Crest3}} was specifically driving high-level GFP expression in transfected somatosensory neurons at all axial levels immediately after NCC emigration and coalescence into DRGs between E2.5 and E4 (Figure 3.3 a-b), in addition to labeling further NCC progeny at early developmental stages (Figure 3.3 c-f). The \textit{IsI1}^{Crest3}-driven GFP expression reflected the endogenous IsI1 protein expression pattern in the sensory lineage, whereas none was found in IsI1 expressing motor neurons or other locations of the neural tube. Fluorescence was also detected in sparse DRG neurons on the non-transfected (contralateral) side, consistent with the findings that NCCs can migrate to either side of the embryo (Figure 3.3 e-f) (Frank and Sanes, 1991). The analysis also verified that the endogenous gene expression pattern can be recapitulated by the use of mouse enhancers in the chick embryo. Additionally, it could have been shown that both orientations of the ECR sequence were capable of modulating gene expression with similar efficiency (data not shown).
3. Results

Figure 3.2: *In silico* ECR screen of the *Islet1* gene locus

(a) Genomic comparison of the mouse *Islet1* (*Isl1*) gene locus between different species reveals evolutionary conserved non-coding regions (ECRs) upstream of the gene transcription start site including the somatosensory-specific enhancer *Isl1Crest3*.

(b) *Isl1Crest3* sequence alignment shows high conservation between human, mouse and chicken, indicated by asterisks.
Due to the stable genomic integration of the ‘enhancer-promoter-reporter gene’ expression cassette, it became possible to overcome the limitations of plasmid-based transient transfection procedures in massively proliferating NCCs and to study the late development of NCC-derived lineages in chick embryos. In accordance with the early
pan-sensory expression pattern, \( Isl1^{Crest3} \) was driving reporter gene expression in both small and large-diameter somatosensory neurons in late-gestation DRGs and visualized the central afferent collateral projections in the spinal cord (Figure 3.4 a-d). Based on the central afferent termination zones in the dorsal and ventral horn of the spinal cord, \( Isl1^{Crest3} \) revealed an unbiased tagging of all three sensory subtypes: nociceptive, mechanoreceptive and proprioceptive neurons.

Consistently, \( Isl1^{Crest3} \) robustly labeled peripheral sensory projections terminating in glabrous skin, feathery skin and musculature (Figure 3.5 a-b). The central connectivity pattern was mirrored by the corresponding sensory end organ structures targeted by their \( Isl1^{Crest3+} \) peripheral collaterals, including nociceptive and mechanoreceptive endings in the dermis (Figure 3.5 c-d) and muscle spindle receptors on intrafusal muscle fibers (Figure 3.5 e-h).
Using this approach, \( Is1^{\text{Crest3+}} \)-tagged central afferent projections of somatosensory neurons were mapped throughout gestation in chick. The \( Is1^{\text{Crest3}} \) enhancer showed robust activity in the somatosensory lineage at least until E15. The labeled developing central afferents entered the spinal cord via the dorsal root entry zone and bifurcated into axon collaterals that extended rostrally and caudally within the dorsal funiculus, prior to penetrating the spinal cord gray matter (Figure 3.6 a-b).
3. Results

The first Isl1<sup>Crest3</sup><sup>+</sup> somatosensory axons entered the spinal cord gray matter at around E7 (Figure 3.7 a-c). These axons were invariably negative for the nociceptive marker TrkA, while the first TrkA<sup>+</sup> axons began advancing into the dorsal horn gray matter between E8 and E9 (Figure 3.7 d-f). These observations were in consistence with previous transganglionic axon tracing experiments in chick (Davis et al., 1989; Eide and Glover, 1997). At around the same developmental stage, the proprioceptive projections began reaching their targets in the intermediate and ventral spinal cord (Figure 3.6 c-d). In late-gestation embryos, Isl1<sup>Crest3</sup>-tagged central afferents converged at their termination zones (Figure 3.6 e-h). Thus, the overall developmental sequence of proprioceptive/mechanoreceptive (TrkA<sup>-</sup>) and nociceptive (TrkA<sup>+</sup>) collateral extension into the gray matter in chick mirrored that observed in the mammalian spinal cord. However, the delay in the establishment of nociceptive
compared to proprioceptive afferent connectivity observed in mammals appeared temporally compressed in avians, likely reflecting an ontogenic adaptation in precocial birds (Fitzgerald, 1987; Mimics and Koerber, 1995; Ozaki and Snider, 1997; Marmigère and Ernfors, 2007).

Figure 3.7: Development of central afferent projections in chick II

(a-f) Genetic tagging of ipsilateral central afferent projections by Isl1\textsuperscript{Crest3} in transverse sections. At E7 the first Isl1\textsuperscript{Crest3}+ TrkA+ axons (closed arrowheads) enter the gray matter of the medial dorsal horn (a-c). First nociceptive projections, labeled by Isl1\textsuperscript{Crest3} and TrkA (open arrowheads), penetrate the gray matter of the lateral dorsal horn at E8 (d-f). Scale bar: 50 µm.

These results therefore establish Isl1\textsuperscript{Crest3} as an early pan-sensory neuron gene regulatory element that can be used for driving a specific and robust reporter gene expression in somatosensory neurons. Furthermore, the analysis confirmed the suitability of the chosen strategy based on stable genetic tagging of DRG neurons in chick by the stable expression system STEVE.
3.3. Analysis of the sensory neuron-specific enhancer \textit{Avil}^Lucy1

Advillin (Avil), a member of the Gelsolin superfamily of actin binding proteins, has been shown to be exclusively expressed in peripheral sensory neurons and has been implicated in regenerative neurite outgrowth (Ravenall \textit{et al.}, 2002; Hasegawa \textit{et al.}, 2007; Lau \textit{et al.}, 2011; Zurborg \textit{et al.}, 2011). Transgenic mice expressing reporter genes under the control of the \textit{Avil} promoter displayed robust and specific expression in postmitotic somatosensory neurons, from embryonic stages into adulthood, thus enabling the study of central and peripheral somatosensory target innervation (Hasegawa \textit{et al.}, 2007; Zurborg \textit{et al.}, 2011). The gene coding sequence of \textit{Avil} showed high homology between mammals, but has not yet been identified in the chick genome. In order to establish a late-gestation and adult pan-sensory marker, the \textit{Avil} gene locus was screened for putative enhancers via \textit{in silico} ECR screen (Figure 3.8 a). An identified ECR, \textit{Avil}^Lucy1, which was located upstream adjacent to the gene coding sequence, was also included in the \textit{Avil} promoter sequences chosen for transgenic mice. This ECR, with a length of 600 bps, showed 80\% homology with the human sequence and included 26 putative conserved TFBS (Figure 3.8 b).

The identified ECR \textit{Avil}^Lucy1 was inserted into the stable expression vector STEVE and analyzed for the spatial and temporal enhancer activity in the chick sensory lineage. \textit{Avil}^Lucy1 was driving a robust reporter gene expression with high specificity in transfected sensory neurons, which retained at least until hatching (Figure 3.9 a-d). Therefore, the \textit{Avil}^Lucy1-mediated expression recapitulated the transgene-driven \textit{Avil} expression pattern observed by other groups (Hasegawa \textit{et al.}, 2007; Zurborg \textit{et al.}, 2011). Compared to \textit{IsI1Crest3}, \textit{Avil}^Lucy1 displayed a later onset of enhancer activity at around E5. By the analysis of peripheral collaterals, \textit{Avil}^Lucy1-tagged projections innervated the dermis (Figure 3.9 e-f) and intrafusal, but not extrafusal muscle fibers (Figure 3.9 g-j). As for \textit{IsI1Crest3}, \textit{Avil}^Lucy1+ central afferent collateral projections were mapped throughout gestation in chick, which closely resembled the observations made by the pan-sensory marker \textit{IsI1Crest3} (Figure 3.10). Herein, \textit{Avil}^Lucy1-labeled central afferents occupied all termination zones in the spinal cord of nociceptive, mechanoreceptive and proprioceptive axons, indicating that \textit{Avil}^Lucy1-driven expression was not biased towards a certain sensory subtype.
3. Results

**Figure 3.8: In silico ECR screen of the Advillin gene locus**

(a) Genomic comparison of the mouse Advillin (Avil) gene locus between different mammalian species reveals an evolutionary conserved non-coding region (ECR), *AvilLucy1*, upstream adjacent to the gene coding sequence.

(b) *AvilLucy1* sequence alignment partially shows high conservation between human, mouse and opossum, indicated by asterisks.

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**Legend:**
- **mouse Advillin locus**
- *AvilLucy1* sequence alignment
- evolutionary conserved non-coding sequence
- evolutionary conserved exon sequence
- evolutionary conserved intron sequence
3. Results

Figure 3.9: Tagging of somatosensory neurons by Avil<sup>*Lucy<sub>1</sub>* in late-gestation chick embryos

(a-d) Avil<sup>*Lucy<sub>1</sub>*-driven reporter gene expression displays specific enhancer activity in somatosensory neurons, in addition to labeling their central afferents in transversal sections of the E9 spinal cord (a-b). Isl1 protein expression is restricted to somatosensory neurons, motor neurons located in the ventral horn and dl3 interneurons. Avil<sup>*Lucy<sub>1</sub>*+ cells overlap with Isl1+ somatosensory neurons (c-d). Scale bar: 200 µm.

(e-f) Avil<sup>*Lucy<sub>1</sub>* labels peripheral somatosensory projections innervating the feathery skin in E12 transverse sections. Nociceptive and mechanoreceptive axons are labeled by βIII-tubulin (Tuj1). Arrowheads indicate feather follicle shafts. Dotted lines demarcate outer side. Scale bar: 100 µm.

(g-j) Avil<sup>*Lucy<sub>1</sub>*+ muscle spindle receptors terminating on intrafusal muscle fibers (closed arrowheads) show no overlap with motor end plates (open arrowheads), detected by Bungarotoxin (Btx), in E12 longitudinal sections (g-h). Tuj1 labels both sensory and motor axons. Detailed view of an innervated muscle spindle (h-j). Scale bar: 100 µm.
These results demonstrate that the newly identified enhancer AvilLucy1 can be used as a pan-sensory neuron marker from mid-embryonic stages at least into late-gestation, by driving specific and strong reporter gene expression in the somatosensory neuron lineage in chick.
3. Results

3.4. Analysis of ipsilaterally and contralaterally migrating neural crest cells

During early vertebrate development, a vast range of peripheral cell types, including autonomic and somatosensory ganglion neurons and further non-neuronal cells, are generated from NCCs, which arise from the roof plate of the dorsal neural tube and emigrate towards their respective peripheral targets (Bronner-Fraser and Fraser, 1988; Frank and Sanes, 1991; Le Douarin and Kalcheim, 1999; Kim et al., 2003; Squire et al., 2008). The adoption of specific cell fates by NCCs is a multistep process involving extensive interaction with the peripheral locales they colonize and is further influenced by the time window of emigration (Harris and Erickson, 2007; Marmigère and Ernfors, 2007). The extent to which the fate of premigratory or early migratory NCCs is predetermined remains less clear, however (Hari et al., 2012). The initial pattern of NCC emigration appears to be determined by homotypic interactions, during which some NCCs enter trajectories that cause them to cross the neural tube midline and to contribute to the contralateral NCC migratory stream (Figure 3.11 a) (Frank and Sanes, 1991; Carmona-Fontaine et al., 2008).

Figure 3.11: Scenarios of ipsilaterally and contralaterally migrating NCC progeny in chick

(a-c) Schematic of a unilaterally transfected neural tube (green label) depicts ipsilateral and contralateral neural crest cell (NCC) streams colonizing dorsal root ganglia (DRGs) (a). Two scenarios of contralaterally migrating NCCs contributing to peripheral neurons and circuits: unbiased, giving rise to all neuron classes (b), or biased to a certain neuron class (c). Numbers: preferential spinal cord laminar targets of nociceptive (TrkA), mechanoreceptive (TrkB) and proprioceptive (TrkC) DRG neuron axon collaterals. S, sympathetic neuron; SCG, sympathetic chain ganglion.
George and colleagues raised the possibility that NCCs diverge into discrete lineage-restricted pools of NCCs choosing diametrically opposed (ipsilateral and contralateral) migratory streams, such that contralateral NCCs are biased to generate pain-sensing nociceptive somatosensory neurons within DRGs (Figure 3.11 c) (George et al., 2007; Lefcort and George, 2007; George et al., 2010). The author’s conclusions were inferred from three principle observations: first, the correlated timing of contralateral NCC and nociceptive DRG neuron differentiation, second, an indirectly estimated bias in the contribution of contralateral NCCs to nociceptive DRG neurons and third, a
3. Results

reduction of nociceptive or total DRG neuron numbers upon surgically cutting off the contralateral NCC stream at the neural tube midline (George et al., 2007).

This hypothesis was investigated by STEVE-based direct lineage-tracking upon unilateral in ovo transfection, followed by neuron type composition analysis in the tagged NCC progeny. Therefore, any lineage-restriction of contralaterally migrating NCCs would consequently be reflected in a bias in the proportion of neuronal classes transfected in contralateral DRGs, compared to those in the ipsilateral DRGs (Figure 3.11 b-c). Upon unilateral neural tube transfection of constitutive (CAG) and somatosensory-specific (Isl1Crest3+) constructs, both ipsilateral and contralateral migrating NCC derivatives were indeed labeled at early and late stages of gestation (Figure 3.12 a-c). Furthermore, the method allowed detailed observation of NCC delamination and migration along distinct pathways (Figure 3.12 d-f). The three major DRG neuron classes form stereotypic laminar connections in the spinal cord via central axon collaterals, reflecting their incorporation into discrete somatosensory circuits (Brown, 1982; Lallemend and Ernfors, 2012). However, in neither the ipsilateral nor the contralateral spinal cord the Isl1Crest3+ central collaterals displayed an obvious bias towards specific termination zones in the dorsal or ventral horn of the spinal cord (Figure 3.12 b-c).

Figure 3.13: Genetic tagging of ipsilateral and contralateral somatosensory neurons in chick

*Figure 3.13: Genetic tagging of ipsilateral and contralateral somatosensory neurons in chick* (a-l) Genetic tagging of ipsilateral and contralateral NCC-derived DRG neurons by somatosensory-specific Isl1Crest3 (a-d) and AvilLucy1 (e-h) and by ubiquitous CAG (i-l) in transverse sections at E9. Isl1, pan-DRG neuron marker. Scale bar: 100 µm.
3. Results

Figure 3.14: DRG neuron type composition of ipsilateral and contralateral NCC progeny in chick

**Figure 3.14** depicts the composition of DRG neurons derived from ipsilateral (a-i) and contralateral (j-r) NCC progenies, tagged by Isl1Crest3. Major DRG neuron classes are highlighted: nociceptive (TrkA), mecanoreceptive (TrkB) and proprioceptive (TrkC) neurons. Co-expression of Isl1 with three independent transgenes: Isl1Crest3, AvilLucy1, or CAG, compared to Isl1+ total DRG neurons. No significant differences in the proportion of eGFP-tagged neurons expressing TrkA, TrkB or TrkC were observed. A substantial portion of DRG neurons express more than one Trk receptor class at E9. Data are presented as mean ± SEM (two-tailed Student’s t test with two samples and unequal variance), ns, not significant, p>0.05; see Table 3.2 for statistical analysis.
3. Results

For a more detailed analysis, the relative proportions of the three major classes of DRG neurons, respectively expressing the Neurotrophin receptors TrkA (corresponding to nociceptive neurons), TrkB (≈low-threshold mechanoreceptive neurons) or TrkC (≈proprioceptive neurons), derived from ipsilaterally versus contralaterally migrating NCCs were compared (Lewin and Moshourab, 2004; Marmigère and Emfors, 2007). Thereby, NCC progeny were labeled upon selectively tracking neuronal or both neuronal and non-neuronal cells with three distinct transgene vectors facilitating DRG neuron-restricted (Isl1\textsuperscript{Crest3}, Avil\textsuperscript{Lucy1}) or generalized (CAG) reporter expression (Figure 3.13). Throughout this analysis, the relative proportions of TrkA\textsuperscript{+}, TrkB\textsuperscript{+} or TrkC\textsuperscript{+} DRG neurons derived from contralateral NCCs (Figure 3.14 j-r) were indistinguishable from those derived from ipsilateral NCCs (Figure 3.14 a-i). The same principal results were consistently obtained by all three constructs (Figure 3.14 s-u). Furthermore, the neuron class composition of both ipsilaterally- and contralaterally-derived NCC progeny matched that characteristically found for Isl1\textsuperscript{+} DRG neurons (Figure 3.14 s-u), together suggesting that contralaterally migrating NCCs lack a measurable bias towards generating particular DRG neuron classes.

In late-gestation embryos, DRG neurons of both ipsilateral and contralateral migratory streams would establish stereotypic laminar connectivity patterns according to their phenotypic profiles. In accordance to the Neurotrophin receptor quantification, the overall connectivity patterns established by contralaterally-derived Isl1\textsuperscript{Crest3+} DRG neurons invariably matched those formed by ipsilaterally-derived Isl1\textsuperscript{Crest3+} DRG neurons, with collaterals projecting to the dorsal horn, intermediate gray matter and the ventral horn of the spinal cord (Figure 3.15 a-b). Contralaterally-derived central afferents were terminating in laminae I-II predominantly targeted by the collaterals of nociceptive DRG neurons, laminae III-V preferentially targeted by low-threshold mechanoreceptive neurons, and laminae VI and IX targeted by proprioceptive neurons (Figure 3.15 c-e) (Brown, 1982; Lallemand and Emfors, 2012).
3. Results

In order to detect a small, but significant laminar bias of the overall connectivity patterns, projection density maps were performed (Figure 3.15 f). Herein, the amount of projections converging onto laminae I-II versus laminae III-V that extended from
ipsilaterally-derived Isl1<sup>Crest3+</sup> DRG neurons (Figure 3.15 g and Figure 3.18), compared to those extending from contralaterally-derived Isl1<sup>Crest3+</sup> neurons (Figure 3.15 h and Figure 3.19) or TrkA<sup>+</sup> nociceptive neurons (Figure 3.15 i and Figure 3.20) was measured. Whereas the vast majority of TrkA<sup>+</sup> collaterals exclusively targeted laminae I-II, both contralateral and ipsilaterally-derived Isl1<sup>Crest3+</sup>-tagged DRG neurons exhibited a slight bias towards laminae III-V (Figure 3.15 j).

In addition, the central connectivity patterns established by ipsilaterally- and contralaterally-derived DRG neurons were mirrored by the corresponding sensory end organ structures targeted by their AvilLucy<sup>L</sup>-tagged peripheral collaterals, including presumptive nociceptive and mechanoreceptive endings in the skin (Figure 3.16 a-d) and muscle spindle receptors on intrafusal muscle fibers (Figure 3.16 e-h). Moreover, ipsilaterally and contralaterally migrating CAG-tagged NCCs contributed to several
other neuronal and non-neuronal NCC derivatives, including Schwann cells (Figure 3.17 a-d), skin melanocytes (Figure 3.17 e-h) and sympathetic ganglion neurons (Figure 3.17 i-l). Ipsilaterally and contralaterally migrating NCCs thus appear to give rise to an equivalent range of neuronal and non-neuronal derivatives.

Using STEVE-based direct lineage-tracking and systematic analysis of NCC progeny and their connections in chick, the results provide conclusive evidence that primary somatosensory neurons derive from neurogenically equivalent ipsilaterally and contralaterally migrating NCCs.
3. Results

Figure 3.18: Projection density map of ipsilaterally-derived $I_{sl^{\text{Crest3+}}}$ central collaterals in chick

(a) Average axon densities were established in serial transverse sections (n=18/8 sections/embryos), normalized and plotted as heat maps (red-blue: high-low axon density).

(b) Heat map of mGFP$^+$ central collaterals labeled by $I_{sl^{\text{Crest3+}}}$ transgene in ipsilateral E12 dorsal horn. Inner and outer dotted lines respectively delineate white (w) and gray (g) matter. Numbers indicate respective laminae. Asterisk, dorsal root entry zone and Lissauer’s tract.

(c) Normalized raw image files of $I_{sl^{\text{Crest3+}}}$-tagged central collaterals in ipsilateral E12 dorsal horn sections (1-18).
3. Results

Figure 3.19: Projection density map of contralaterally-derived \( \text{Isla}^{\text{Crest3}} \) central collaterals in chick

(a) Average axon densities were established in serial transverse sections (n=18/8 sections/embryos), normalized and plotted as heat maps (red-blue: high-low axon density).

(b) Heat map of mGFP\(^+\) central collaterals labeled by \( \text{Isla}^{\text{Crest3}} \) transgene in contralateral E12 dorsal horn. Inner and outer dotted lines respectively delineate white (w) and gray (g) matter. Numbers indicate respective laminae. Asterisk, dorsal root entry zone and Lissauer's tract.

(c) Normalized raw image files of \( \text{Isla}^{\text{Crest3}} \)-tagged central collaterals in contralateral E12 dorsal horn sections (1-18).
3. Results

**Figure 3.20: Projection density map of TrkA⁺ central collaterals in chick**

(a) Average axon densities were established in serial transverse sections (n=18/8 sections/embryos), normalized and plotted as heat maps (red-blue: high-low axon density).

(b) Heat map of TrkA⁺ central collaterals in E12 dorsal horn. Inner and outer dotted lines respectively delineate white (w) and gray (g) matter. Numbers indicate respective laminae. Asterisk, dorsal root entry zone and Lissauer's tract.

(c) Normalized raw image files of TrkA⁺ central collaterals in E12 dorsal horn sections (1-18).
### 3. Results

#### Table 3.2: Statistical analysis of ipsilateral and contralateral NCC progeny experiments

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<tr>
<td>CAG contra</td>
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**Data of colocalization and projection density analysis are presented as the mean ± standard error of the mean (SEM). Statistical comparisons were established using a two-tailed Student's t test with two samples and unequal variance.**
3.5. Analysis of the mechanoreceptor-specific enhancer $\text{Ntng1}^{\text{Mech1}}$

By exploring recent literature and public gene expression data bases for mRNAs, one of the identified candidates, NetrinG1 ($\text{Ntng1}$), appeared to be expressed by a subset of large-diameter somatosensory neurons in posnatal mouse DRGs (Yin et al., 2002; www.brain-map.org). $\text{Ntng1}$ encodes a GPI-linked synaptic adhesion molecule and its selective interaction with the transmembrane NetrinG1 ligand (NGL1) is implicated in axonal outgrowth, lamina/pathway-specific differentiation of dendrites and synapse formation, whereas its deregulation is associated with diverse brain dysfunctions (Nakashiba et al., 2002; Nishimura-Akiyoshi et al., 2007; Woo et al., 2009).

**Figure 3.21: In silico ECR screen of the NetrinG1 gene locus**
(a) Genomic comparison of the chicken NetrinG1 ($\text{Ntng1}$) gene locus between different species reveals an evolutionary conserved non-coding region (ECR), $\text{Ntng1}^{\text{Mech1}}$, 2.6 kb upstream of the gene transcription start site.
(b) $\text{Ntng1}^{\text{Mech1}}$ sequence alignment shows high conservation between human, mouse and chicken, indicated by asterisks.
Although the role of Ntng1 in sensory neurons remains unknown, one might speculate, based on previous observations in the brain, that Ntng1-NGL1 interactions could also have an impact on the establishment of somatosensory circuits (Woo et al., 2009). The Ntng1 gene locus was screened via in silico ECR analysis for a putative enhancer driving the sensory subtype-specific gene expression (Figure 3.21 a). One of the ECRs identified, Ntng1 Mech1, located 2.6 kb upstream of the gene coding sequence in the chick, showed 86% and 83% homologies with the human and mouse sequences, respectively, and included 26 putative TFBS (Figure 3.21 b). In the human and the mouse genome the evolutionary conserved Ntng1 Mech1 was detected in the first intron region.

In order to analyze the potential enhancer activity of Ntng1 Mech1, the ECR was coupled to STEVE and tested for the spatial and temporal expression pattern in both early and late stages of the somatosensory lineage. Ntng1 Mech1 consistently labeled a population of large-diameter sensory neurons confined to the ventrolateral portion of the DRGs (Figure 3.22 a-b), in contrast to the pan-sensory reporter gene expression driven by Isl1 Crest3 (Figure 3.22 c-d). Thereby, Ntng1 Mech1 appeared to reflect the endogenous Ntng1 expression pattern in somatosensory neurons observed by other groups (Yin et al., 2002; www.brain-map.org). The earliest activity of Ntng1 Mech1 in somatosensory neurons could be detected after E6, while high-level expression persisted from E9 to at least until hatching. An additional activity was observed in a small subset of ventral interneurons in the spinal cord after E5. The central afferents of Ntng1 Mech1-labeled sensory neurons displayed a highly restricted projection pattern confined to the medial dorsal horn of the spinal cord (Figure 3.22 e-f), in contrast to Isl1 Crest3+ central collaterals occupying all laminae of the dorsal horn, as well as the intermediate and ventral spinal cord (Figure 3.22 g-h). At E7, the first detectable Ntng1 Mech1+ centrally projecting axons assumed a medial position within the dorsal funiculus and segregated from laterally positioned TrkA+ nociceptive projections prior to entering the spinal cord gray matter (compare Figure 3.23 a-b and i-j). By E9, Ntng1 Mech1+ axons penetrated the dorsal gray matter, but remained confined within the medioventral quadrant of the dorsal horn, while TrkA+ nociceptive afferents began occupying most of the lateral portion of the dorsal horn gray matter (Figure 3.23 c-d). At the same time, Isl1 Crest3+ nociceptive and mechanoreceptive afferents projected throughout the dorsal horn,
whereas \textit{Isl1}^{\text{Crest3}}\textsuperscript{-}labeled proprioceptive collaterals occupied the intermediate and ventral spinal cord (Figure 3.23 k-l and Figure 3.6 c-d).

**Figure 3.22: Selective tagging of a somatosensory neuron subtype by \textit{Ntng1}^{\text{Mech1}} in chick**

(a-d) \textit{Ntng1}^{\text{Mech1}} enhancer activity selectively drives reporter gene expression in large-diameter ventrolateral dorsal root ganglion (DRG) neurons (a-b), compared to \textit{Isl1}^{\text{Crest3}}-driven GFP expression in all somatosensory neuron types (c-d) in transverse sections at E9. Isl1, pan-DRG neuron marker. Scale bar: 100 µm.

(e-g) \textit{Ntng1}^{\text{Mech1}}+ central collaterals project to the medioventral dorsal horn in transverse sections at E12 (e-f). Additional low \textit{Ntng1}^{\text{Mech1}} activity is detected in ventral interneurons (asterisks). In contrast, \textit{Isl1}^{\text{Crest3}}+ central collaterals occupy all laminae of the dorsal horn, as well as the intermediate and ventral spinal cord (g-h). NeuN\textsuperscript{+} neurons outline spinal cord gray matter. Numbers indicate respective laminae. DH, dorsal horn; VH, ventral horn. Scale bar: 100 µm.

Between E12 and E15, \textit{Ntng1}^{\text{Mech1}}+ axons converged onto an increasingly condensed zone within the medioventral quadrant of the dorsal horn, which remained largely unoccupied by TrkA\textsuperscript{+} nociceptive projections (compare Figure 3.23 e-h and m-p). The identified \textit{Ntng1}^{\text{Mech1}} thus has been shown to drive specific expression in a subset of somatosensory neurons, whose central collaterals occupied a central termination zone distinct from that occupied by nociceptive or proprioceptive afferents. Based on their large soma size, their defined ventrolateral position in the DRG, and their discrete
central afferent termination pattern in the dorsal horn, Ntng1^{Mech1+} neurons showed initial characteristics of low-threshold mechanoreceptors (LTMRs).

For a more detailed analysis, the relative proportions of Neurotrophin receptors expressing neurons tagged by Ntng1^{Mech1+} compared to Isl1^{Crest3+}-tagged neurons were scored. Herein, the three major classes of DRG neurons expressing TrkA, TrkB and TrkC mainly correspond to nociceptive neurons, subtypes of LTMR neurons and proprioceptive neurons, respectively (Figure 3.24 a-i) (Lewin and Moshourab, 2004; Marmigère and Ernfors, 2007).
Around 78.05 ± 2.68 % of Isl1\textsuperscript{Crest3}-tagged DRG neurons were TrkA\textsuperscript{+} (Figure 3.24p), which matches the proportion of TrkA\textsuperscript{+} neurons in total Isl1\textsuperscript{+} DRG neurons (Figure 3.14s), and that typically found in the DRGs of adult mammals (Fariñas et al., 1998; Marmigère and Ernfors, 2007). In contrast, only 9.44 ± 2.67 % of Ntng1\textsuperscript{Mech1+}...
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DRG neurons expressed the nociceptive marker TrkA (Figure 3.24 p). At the same time, Ntng1\textsuperscript{Mech1+} neurons contained markedly higher proportions of TrkB\textsuperscript{+} and TrkC\textsuperscript{+} neurons with 33.24 ± 3.45 % and 27.98 ± 5.22 %, respectively, compared to Isl1\textsuperscript{Crest3+}-tagged sensory neurons with 19.80 ± 1.21 % and 19.07 ± 1.78 %, respectively (Figure 3.24 p). In agreement with previous data, the cumulative proportion of TrkA\textsuperscript{+}, TrkB\textsuperscript{+} and TrkC\textsuperscript{+} neurons scored in total or Isl1\textsuperscript{Crest3+} DRG neurons amounted to more than 100%, reflecting the presence of sensory neurons co-expressing more than one class of Trk receptors (Fariñas et al., 1998). Taking this into account, a considerable portion of Ntng1\textsuperscript{Mech1+} neurons appeared to express neither Trk receptor class, which would be in congruence with the molecularly identified earlyRet\textsuperscript{+} population of LTMR neurons in mouse (Bourane et al., 2009; Luo et al., 2009). Therefore, the Ntng1\textsuperscript{Mech1+}-tagged DRG neurons were analyzed for co-expression with the myelination marker NF200, as well as with Ret, a newly identified marker predominantly labeling rapidly adapting LTMRs and subtypes of nociceptive neurons (Figure 3.24 j-o) (Bourane et al., 2009; Luo et al., 2009). The vast majority of Ntng1\textsuperscript{Mech1+} sensory neurons were NF200\textsuperscript{+} (96.42 ± 1.35 %), whereas only 31.04 ± 3.95 % co-expressed Ret (Figure 3.24 q). According to the obtained molecular marker profile, Ntng1\textsuperscript{Mech1+} neurons thus partially separate from Ret\textsuperscript{+} and Trk\textsuperscript{+} mechanoreceptive populations, and appear to selectively tag a molecularly not-yet-identified subtype of Aβ-LTMRs.

Since the specification of sensory neuron classes is associated with characteristic differences in their cell soma sizes, the distribution of soma sizes of Ntng1\textsuperscript{Mech1+} neurons, compared to Isl1\textsuperscript{Crest3+} neurons, as well as those of TrkA\textsuperscript{+}, TrkB\textsuperscript{+} and TrkC\textsuperscript{+} neurons were measured (Figure 3.25). Ntng1\textsuperscript{Mech1+} neurons showed a marked bias towards the largest DRG neuron sizes with an average soma size area of 309.09 ± 2.72 µm\textsuperscript{2} (Figure 3.25 a, f). Furthermore, the distribution of Ntng1\textsuperscript{Mech1+} cell sizes clearly segregated from the overall distribution of Isl1\textsuperscript{Crest3+} soma sizes with an average of 188.27 ± 1.98 µm\textsuperscript{2} (Figure 3.25 b, f), and from that of the small-diameter TrkA\textsuperscript{+} neurons with an average of 138.43 ± 1.65 µm\textsuperscript{2} (Figure 3.25 c, f). In contrast, Ntng1\textsuperscript{Mech1+} neuron soma sizes partially overlapped with those of TrkB\textsuperscript{+} and TrkC\textsuperscript{+} neurons with 264.54 ± 3.24 µm\textsuperscript{2} and 251.54 ± 3.36 µm\textsuperscript{2}, respectively (Figure 3.25 d-f). The large soma size of DRG neurons tagged by Ntng1\textsuperscript{Mech1} was thus in congruence with their Aβ-LTMR-specific molecular marker profile.
3. Results

Both adult and embryonic nociceptive and mechanoreceptive sensory neurons can be distinguished by their electrophysiological properties, including characteristic action potential waveforms (Koerber et al., 1988; Djouhri et al., 1998; Fang et al., 2005). Mechanoreceptors typically fire narrow uninflected spikes with short half peak durations (HPDs) and short afterhyperpolarization (AHP) durations (Figure 3.26 a), whereas nociceptors fire action potentials that exhibit a 'hump' on the falling phase.
and have very long HPDs and AHP durations (Figure 3.26 b) (Lechner et al., 2009). Via whole cell patch clamp recordings in dissociated E9 DRG neurons, which were performed in collaboration with Stefan G. Lechner and Gary R. Lewin at the ‘Max-Delbrück-Center for Molecular Medicine’ in Berlin, the firing properties of Ntng1 Mech1+ compared to Isl1 Crest3+ sensory neurons were monitored. Action potentials evoked by current injection of patched Ntng1 Mech1+ neurons displayed short HPDs (1.55 ± 0.11 ms) and AHPs (3.92 ± 0.25 ms) (Figure 3.26 e-f, h), which were characteristic for mechanoreceptors. Isl1 Crest3+ neurons, marking all sensory neuron subtypes, displayed a considerably wider range of HPD and AHP values, including both nociceptor- and mechanoreceptor-specific types of action potentials (Figure 3.26 e-f). Herein, the average HPD of 2.50 ± 0.18 ms and average AHP duration of 6.74 ± 0.84 ms evoked from Isl1 Crest3+ neurons was significantly higher than from Ntng1 Mech1+-tagged neurons (Figure 3.26 h). Ntng1 Mech1+ neurons therefore represent a discrete population of sensory neurons with spike characteristics typical of embryonic mechanoreceptors.

LTMRs can be further distinguished by the feature of a rapid and highly sensitive mechanosensitive current, which appears as soon as they innervate their peripheral targets (Lechner et al., 2009). Adult and embryonic LTMRs possess a so-called RA-mechanosensitive current, which is a mechanically activated inward current that activates and inactivates within a few milliseconds (Figure 3.26 c) (Drew et al., 2002; Hu and Lewin, 2006; Lechner et al., 2009). RA-mechanosensitive currents are also found in nociceptors, but the majority of nociceptors exhibit much slower inactivating IA and SA-currents (Figure 3.26 d). By investigating mechanosensitive currents Ntng1 Mech1+ neurons exclusively displayed RA-currents with an average inactivation time of 1.28 ± 0.32 ms, whereas Isl1 Crest3+ neurons exhibited both RA- and IA-currents with an average inactivation time of 7.36 ± 1.46 ms (Figure 3.26 g-h). Consistent with previous observations showing that SA-currents appear very late in mouse DRG, SA-currents were absent from chick E9 nociceptors (Lechner et al., 2009).
For electrophysiological measurements, Ntng1^{Mech1+} and Isl1^{Crest3+} neurons spanning the whole range of their cell size distribution were selected (compare Figure 3.27 and
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Figure 3.25 a-b). Ntng1\textsuperscript{Mech1+} neurons thus represent a discrete population of somatosensory neurons with functional properties characteristic for LTMRs, including narrow uninflected action potentials with short HPDs, short AHPs and RA-mechanosensitive currents.

In mammals, the majority of nociceptive afferents connect to relay neurons within the most superficial laminae I and II of the dorsal horn, while LTMRs mainly terminate throughout the successively deeper laminae II (inner) and V, and proprioceptive afferents extend their axons in laminae VI and XI of the intermediate and ventrolateral gray matter (Brown, 1982; Lallemend and Ernfors, 2012). Unlike in mammals, the dorsal horn in chick displays a mediolaterally rotated appearance, with neurons expressing the laminae I-III marker Lmx1b respectively clustering into a lateral (laminae I-II) and a smaller mediodorsal crescent-shaped domain (lamina III) in the dorsal horn (Figure 3.28 a-b) (Rebelo et al., 2010; Wild et al., 2010). In order to obtain more precise information, the stereotypic laminar connectivity pattern established by Ntng1\textsuperscript{Mech1+} DRG neurons was compared with Isl1\textsuperscript{Crest3+} and TrkA\textsuperscript{*} central afferent projections. The vast majority of TrkA\textsuperscript{*} nociceptive projections into the dorsal horn gray matter concentrated within the lateral Lmx1b\textsuperscript{*} domain, while excluding the mediodorsal Lmx1b\textsuperscript{*} crescent (Figure 3.28 g-h), suggesting that the latter corresponds to lamina III, while the former corresponds to laminae I-II (Eide and Glover, 1997; Wild et al., 2010).
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**Figure 3.28:** Central connectivity pattern of $\text{Ntng}^{\text{Mech1}+}$ somatosensory neurons in chick

(a-h) Central collateral projections established by $\text{Ntng}^{\text{Mech1}+}$, $\text{Isl1}^{\text{Crest3}+}$ and $\text{TrkA}^+$ DRG neurons in transverse dorsal horn sections at E12. The discrete laminae in the chick dorsal horn can be distinguished by $\text{Lmx1b}$ which labels laminae I-II on the lateral site and lamina III on the mediodorsal site (a-b). $\text{Ntng}^{\text{Mech1}+}$ afferents terminate in a definite domain ventrally of lamina III, presumably in laminae IV-V (c-d), compared to $\text{Isl1}^{\text{Crest3}+}$ projections occupying all termination zones in the dorsal horn (e-f). Nociceptive $\text{TrkA}^+$ projections converge in laminae I-II (g-h). Neun labels all neurons of the dorsal horn gray matter. Note: compared to mammals, dorsal horn lamination in chick exhibits a mediolaterally rotated appearance. Numbers indicate respective laminae. Dotted lines demarcate white (w) and gray (g) matter. Asterisks, dorsal root entry zone and Lissauer's tract. Scale bar: 100 µm.

(i-l) Comparative projection density maps of $\text{Ntng}^{\text{Mech1}+}$ (g), $\text{Isl1}^{\text{Crest3}+}$ (k) or $\text{TrkA}^+$ nociceptive (l) axon collateral projections in the E12 dorsal horn. Average axon densities were established in serial transverse sections (n=18/8 sections/embryos), normalized and plotted as heat maps (red-blue: high-low axon density) (i). $\text{Ntng}^{\text{Mech1}+}$ projections (j) converge within a medioventral quadrant of the dorsal horn (laminae IV-V), whereas $\text{Isl1}^{\text{Crest3}+}$ projections (k) are distributed among all laminae and $\text{TrkA}^+$ projections (l) on laminae I-II. See Figure 3.18, 3.20 and 3.29 for details on projection density map compositions.

(m) Comparative quantitative analysis of axon densities in laminae I-II versus laminae III-V. Marked bias of $\text{Ntng}^{\text{Mech1}+}$ axons towards laminae III-V, in contrast to slight bias of $\text{Isl1}^{\text{Crest3}+}$ axons towards laminae III-V and marked bias of $\text{TrkA}^+$ projections towards laminae I-II. Data are presented as mean ± SEM (two-tailed Student's $t$ test with two samples and unequal variance). ***, $p<0.001$; see Table 3.4 for statistical analysis.
In accordance to previous obtained observations, Isl1<sup>Crest3</sup>+ axons projected throughout the dorsal horn (Figure 3.28 e-f), while Ntng1<sup>Mech1</sup>+ afferents converged within an oval shaped domain in the medioventral quadrant of the dorsal horn that excluded the mediodorsal Lmx1b<sup>*</sup> crescent (Figure 3.28 c-d). Therefore, Ntng1<sup>Mech1</sup>+ afferents are suggested to terminate mainly in laminae IV-V, as there are currently no markers available in chick to discriminate these laminae further. The significant laminar bias towards laminae IV-V of Ntng1<sup>Mech1</sup>-tagged central collaterals was further confirmed by projection density maps of the overall connectivity pattern (Figure 3.28 j, m and Figure 3.29). In contrast, only a slight bias of Isl1<sup>Crest3</sup>+ axons towards laminae III-V (Figure 3.28 k, m and Figure 3.18), and a marked bias of TrkA<sup>*</sup> nociceptive projections towards laminae I-II was detected (Figure 3.28 l-m and Figure 3.20). Thus, Ntng1<sup>Mech1</sup> defines a novel subtype of LTMRs whose central collaterals exclude laminae III and selectively terminate within a medioventral domain of the dorsal horn presumably corresponding to laminae IV-V.

The in silico-to-in vivo ECR screen in combination with the stable expression system STEVE facilitates identification and comprehensive analysis of late-onset somatosensory-specific genetic markers inaccessible to previous methods. Thereby, a novel molecularly defined subset of somatosensory neurons labeled by Ntng1<sup>Mech1</sup> was uncovered, whose molecular (Trk, Ret, NF200 expression), morphological (soma size), electrophysiological (action potential waveform), functional (mechanotransduction current) and anatomical (central connectivity) characteristics correspond to a discrete subclass of Aβ-LTMRs.
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**Figure 3.29: Projection density map of Ntng$^{\text{Mech}1+}$ central collaterals in chick**

(a) Average axon densities were established in serial transverse sections (n=18/8 sections/embryos), normalized and plotted as heat maps (red-blue: high-low axon density).

(b) Heat map of mGFP$^+$ central collaterals labeled by Ntng$^{\text{Mech}1+}$ transgene in E12 dorsal horn. Inner and outer dotted lines respectively delineate white (w) and gray (g) matter. Numbers indicate respective laminae. Asterisk, dorsal root entry zone and Lissauer's tract.

(c) Normalized raw image files of Ntng$^{\text{Mech}1+}$-tagged central collaterals in E12 dorsal horn sections (1-18).
## 3. Results

### Table 3.3: Statistical analysis of Ntng1\(^{\text{Mech}}\) colocalization and cell size experiments

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<td></td>
<td>mean [µm²]</td>
<td>SEM [µm²]</td>
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#### TrkA colocalization / section

| Ntng1\(^{\text{Mech}}\) | 9.44 | 2.67 | 19 | 192 | 1.33 \(\times\) 10\(^{-4}\) |
| Isf1\(^{\text{CREM}}\)    | 78.05 | 2.66 | 15 | 959 | |

#### TrkB colocalization / section

| Ntng1\(^{\text{Mech}}\) | 33.24 | 3.45 | 40 | 340 | 0.0006 |
| Isf1\(^{\text{CREM}}\)    | 19.80 | 1.21 | 19 | 898 | |

#### TrkC colocalization / section

| Ntng1\(^{\text{Mech}}\) | 27.98 | 5.22 | 19 | 178 | 0.1208 |
| Isf1\(^{\text{CREM}}\)    | 19.07 | 1.78 | 13 | 502 | |

#### NF200 colocalization / section

| Ntng1\(^{\text{Mech}}\) | 95.42 | 1.35 | 11 | 276 | |

#### Ret colocalization / section

| Ntng1\(^{\text{Mech}}\) | 31.04 | 3.95 | 11 | 276 | |

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Data of colocalization and cell size analysis are presented as the mean ± standard error of the mean (SEM). Statistical comparisons were established using a two-tailed Student’s t test with two samples and unequal variance.
3. Results

Table 3.4: Statistical analysis of \textit{Ntng}^{1\text{Mech}} electrophysiology and projection density experiments

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Table 3.4: Statistical analysis of \textit{Ntng}^{1\text{Mech}} electrophysiology and projection density experiments

Data of electrophysiology and projection density analysis are presented as the mean ± standard error of the mean (SEM). Statistical comparisons were established using a two-tailed Student’s \textit{t} test with two samples and unequal variance.
3.6. Overall organization of somatosensory central collaterals in chick

The combination of STEVE-mediated somatosensory-specific genetic markers with unbiased tagging of contralateral somatosensory neurons affords selective tracing of central afferent projections throughout the whole embryonic development, unobstructed by additional sites of labeling in the spinal cord. A further benefit is the comparatively sparse axonal labeling originating from small numbers of contralaterally transfected sensory neurons, which can be tuned to allow visualization of individual axon collaterals – a highly useful feature for the development of single neuron connectivity (Chen AI et al., 2006; Jefferis and Livet, 2012). In order to establish the overall central connectivity pattern in chick, trajectories of individual afferent collaterals originating from sparsely labeled Isl1Crest3+, AvilLucy1+ and Ntng1Mech1+ contralateral somatosensory neurons were traced and analyzed in detail.

Central afferents were categorized as following: nociceptive (Isl1Crest3+ TrkA+ collaterals), mechanoreceptive (Isl1Crest3+ TrkA− collaterals terminating in the dorsal horn), ‘laminae IV-V’ mechanoreceptive (Ntng1Mech1+ collaterals) and proprioceptive (Isl1Crest3+ TrkA− collaterals terminating in the intermediate and ventrolateral gray matter) (Figure 3.30 a-f). In agreement with previous observations, the dorsal horn in chick displays a mediolateral rotation, compared to the mammalian dorsal horn organization (Figure 3.30 g-h) (Brown, 1982; Rebelo et al., 2010; Wild et al., 2010). The majority of nociceptive collaterals appeared to directly terminate within laminae I-II upon entering the gray matter via the lateral funiculus/Lissauer’s tract (Figure 3.30 a and Figure 3.31 e-g). In addition, subsets of TrkA+ afferents projected medially along the ventral border of the dorsal horn before veering dorsally along the medial border of laminae I-II, or further medially towards the dorsal funiculus (Figure 3.30 b and Figure 3.31 h-j). This consistent observation apparently revealed a part of a yet uncharted ‘nociceptive tract’ (Figure 3.31 a-d) (Davis et al., 1989; Guo et al., 2011). Both mechanoreceptive and proprioceptive projections entered the medial dorsal horn through the dorsal funiculus, while avoiding laminae I-II (Figure 3.30 c-f). Thereby, most of mechanoreceptor collaterals directly terminated in laminae III-V proximal to their funicular point of entry (Figure 3.30 c and Figure 3.32 a-c).
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Consistently, the majority of ‘laminae IV-V’ mechanoreceptor afferents directly projected to their termination zone ventral to lamina III in the dorsal horn, while collaterals branching to both laminae III and IV-V was rarely observed (Figure 3.30 d and Figure 3.32 d-f).
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Figure 3.31: Trajectory tracings of nociceptive central afferents in chick

(a-d) TrkA⁺ nociceptive axons enter the gray matter of the dorsal horn via the lateral funiculus (LF) and target directly within laminae I-II in transverse dorsal horn sections at E12 and E15. A subset of these collaterals project medially along the ventral border of the dorsal horn (arrowheads) (a), or further towards the dorsal funiculus (DF), as part of a ‘nociceptive tract’ (arrowheads) (b-c). A summarized model of nociceptive afferents is shown in (d). Inner and outer dotted lines respectively delineate white and gray matter. Numbers indicate respective laminae. LT, Lissauer’s tract; GB, glycogen body; CC, central canal; interrogation mark, unknown termination. Scale bar: 100 µm.

(e-g) Example of an individual nociceptive afferent by contralaterally-derived Isl1Crest3⁺ sensory neurons in transverse E12 dorsal horn sections (e), a trajectory reconstruction of different sections (f) and a summarized model of nociceptive afferents entering the dorsal horn via the LF and terminating in laminae I-II (g). Dotted lines differentiate white and gray matter. Scale bar: 100 µm.

(h-j) Example of an individual nociceptive afferent by contralaterally-derived Isl1Crest3⁺ sensory neurons in transverse E12 dorsal horn sections (h), a trajectory reconstruction of different sections (i) and a summarized model of nociceptive afferents projecting from the LF along the ‘nociceptive tract’ (j). Scale bar: 100 µm.
Finally, proprioceptive collaterals transited the gray matter by projecting throughout the medial half of the dorsal horn, while neither type Ia nor type Ib/Ila afferents show differences regarding the position of their funicular entry points, or the angle of their initial trajectories (Figure 3.30 e-f and Figure 3.33 a-f). Furthermore, in analogy to previous observations, proprioceptive afferents showed an extensive axon branching along their trajectories before establishing their final connectivity in laminae VI and IX (Figure 3.33 g-n) (Chen AI et al., 2006). Additionally, \textit{Isl}^{\text{Crest3}-}\text{ and } \textit{Avil}^{\text{Lucy1}-}\text{tagged contralaterally-derived central afferent projections were mapped throughout gestation and reflected the established developmental sequence of somatosensory collateral extension into the spinal cord gray matter (Figure 3.34).

These observations thus support the long-standing idea that afferent connectivity in the spinal cord is established by a majority of central afferent collaterals that directly home in on their respective target zones (Figure 3.30 i) (Fitzgerald, 1987; Davis \textit{et al.}, 1989; Ozaki and Snider, 1997; Redmond \textit{et al.}, 1997; Fitzgerald, 2005).
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Figure 3.33: Trajectory tracings of proprioceptive central afferents in chick

(a-c) Example of an individual proprioceptive afferent by contralaterally-derived Isl1\textsuperscript{Crest3+} sensory neurons in transverse E12 dorsal horn sections (a), a trajectory reconstruction of different sections (b) and a summarized model of proprioceptive afferents projecting throughout the dorsal funiculus (DF) only to lamina VI (c). Dotted lines differentiate white and gray matter. Numbers indicate respective laminae. LF, lateral funiculus. Scale bar: 100 µm.

(d-f) Example of an individual proprioceptive afferent by contralaterally-derived Isl1\textsuperscript{Crest3+} sensory neurons in transverse E12 dorsal horn sections (d), a trajectory reconstruction of different sections (e) and a summarized model of proprioceptive afferents projecting throughout the DF to lamina VI and lamina IX (f). Scale bar: 100 µm.

(g-n) Sparse central afferent axon labeling by contralaterally-derived Isl1\textsuperscript{Crest3+}-labeled sensory neurons in transverse E12 spinal cord sections facilitates delineation of single axon projections (g-j). Detailed view of proprioceptive afferents (outlined in green) shows an extensive axon branching along their trajectories (arrowheads) (k-n). Scale bar: 100 µm.
Figure 3.34: Tagging of contralateral central collaterals throughout gestation in chick

(a-p) Genetic tagging of contralateral central afferent projections by Isl1Crest3 (a-h) and AvilLucy1 (i-p) throughout gestation in transverse sections from E7 to E15. Proprioceptive afferents are indicated by arrowheads, nociceptive sensory axons are labeled by TrkA, motor neurons (MN) and dorsal interneurons (di3 IN) by Isl1. Asterisks, mGFP+ ipsilateral commissural projections. Scale bar: 100 µm.
3. Results

3.7. Peripheral connectivity of Ntng1\(^{\text{Mech}^+}\) somatosensory neurons

The topographic arrangement of sensory subtype-specific central connectivity in the dorsal horn of the spinal cord is expected to be reflected by the innervation of corresponding sensory end organs in defined regions in the periphery (Arber, 2012). Different types of functionally specialized LTMRs along with their corresponding receptor end organs in the skin encode the different qualities of touch (Li et al., 2011).

In mammals, rapidly adapting (RA) A\(\beta\)-LTMRs innervate hair follicles, Meissner corpuscles and Pacinian corpuscles, that selectively respond to hair deflection and to low and high-frequency vibration, respectively (Lewin and Moshourab, 2004; Lumpkin et al., 2010). Slowly adapting (SA) A\(\beta\)-LTMRs terminate in Merkel discs and Ruffini corpuscles, which relay static responses to dermal stretch and indentation (Lewin and Moshourab, 2004; Lumpkin et al., 2010). Pacinian corpuscles and Merkel cells are present in both glabrous and hairy skin. In contrast, Meissner corpuscles and Ruffini corpuscles are exclusively found in glabrous skin, whereas hair follicles are restricted to hairy skin. It is therefore possible to differentiate LTMRs due to their peripheral innervation of distinct morphologically unique end organs and their characteristic physiological properties. Although sensory end organs are well documented in mammals, there is only limited information on avian mechanoperception. In birds, two main types of mechanoreceptors transduce information: Herbst and Merkel corpuscles (Gottschaldt 1985; Necker 2000). The most common and widely distributed Herbst corpuscle, which is considered to resemble the Pacinian corpuscle of mammals, is sensitive to vibration and acceleration (Gottschaldt, 1985; Necker, 2000). Herbst corpuscles are thought to form terminal arborizations in the collar region of most feather follicles and to terminate in the dermis of glabrous skin in digits (Duc et al., 1993). In contrast, Merkel cell receptors, which respond to pressure, are primarily found in featherless skin of avians and differ from Merkel cell-neurite complexes in mammals by accumulating and forming rudimentary Meissner-like corpuscles (Winkelmann and Myers, 1961; Duc et al., 1993; Necker, 2000; Halata et al., 2003). Unlike in mammals, Merkel corpuscles in chick are exclusively found in the superficial layer of the dermis, while the epidermis does not contain any nerve endings (Halata et al., 2003). On the basis of electrophysiological studies, the literature is so far not
conclusive in ranking Herbst and Merkel corpuscles among defined RA or SA modalities (Necker, 1985; Gentle, 1989; Necker, 1990; Duc et al., 1993).

Figure 3.35: Analysis of Ntng1Mech1+ peripheral collaterals in chick glabrous skin

(a-d) Genetic tagging of peripheral afferent projections by Ntng1Mech1+ and AvilLucy1 in digits at E15. Ntng1Mech1+ peripheral somatosensory afferents display a subtype-specific innervation of glabrous skin in the digit (a-b), in contrast to highly complex innervation of AvilLucy1+ somatosensory neurons including all mechanoreceptive and nociceptive subtypes (c-d). Neurofilament (NF) labels total sensory projections. Scale bar: 200 µm.

Ntng1Mech1+ appears to define a subset of LTMRs, whose central connectivity is restricted to laminae IV-V of the dorsal horn. In order to determine whether Ntng1Mech1+ and Ntng1Mech1- neurons are tuned to specific mechanosensory submodalities and terminate in different sensory end organs in the dermis, the peripheral innervation of Ntng1Mech1+ DRG neurons was analyzed. In consistence with labeling central collaterals, Ntng1Mech1+-tagged peripheral projections were observed at different developmental stages at least until hatching. At E15, Ntng1Mech1+ peripheral sensory afferents displayed a subtype-specific innervation of the glabrous skin in digits (Figure 3.35 a-b), in contrast to the widely distributed and highly complex innervation of robustly labeled sensory projections by the pan-somatosensory marker AvilLucy1.
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(Figure 3.35 c-d). No specific innervation of Ntng1\(^{\text{Mech1+}}\) LTMRs was detected in feathery skin of the hindlimb or the dorsal trunk (Figure 3.36 a-b), whereas \(\text{Isl1}^{\text{Crest3}}\)-tagged peripheral afferents were highly accumulated in collar regions of feather follicles and displayed a typical sensory innervation of feathery skin (Figure 3.36 c-d). Since \(\text{Ntng1}^{\text{Mech1}}\) did not drive expression in proprioreceptive neurons, no musculature innervation of Ntng1\(^{\text{Mech1+}}\) peripheral collaterals could be observed.

Both of the main mechanoreceptor end organs, Herbst and Merkel corpuscles, are present in chick glabrous skin. In order to obtain more precise information, whether Ntng1\(^{\text{Mech1+}}\) DRG neurons terminate in a discrete sensory end organ, the peripheral innervation was analyzed morphologically in more detail. At E12, Ntng1\(^{\text{Mech1+}}\) peripheral endings were visualized in the plantar and scaly glabrous skin of the digits,

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**Figure 3.36: Analysis of Ntng1\(^{\text{Mech1+}}\) peripheral collaterals in chick feathery skin**

(a-d) Genetic tagging of peripheral afferent projections by Ntng1\(^{\text{Mech1}}\) and Isl1\(^{\text{Crest3}}\) in the dorsal trunk at E15. No specific innervation of Ntng1\(^{\text{Mech1+}}\) peripheral afferents is detected in feathery skin (a-b). Sparse non-specific mGFP expression is detected in limited regions (asterisks). Peripheral projections labeled by pan-somatosen-sory marker Isl1\(^{\text{Crest3}}\) highly accumulate in collar regions of feather follicles (dotted lines) (c-d). Isl1\(^{\text{Crest3+}}\) axons show complete overlap with neurofilament (NF) illustrating a typical sensory skin innervation. Scale bar: 200 µm.
but no obvious end organ structures could be identified (Figure 3.37). At the latest analyzed embryonic stage, E19, $\text{Ntng}^{\text{Mech1}^+}$-tagged peripheral afferents were observed to project in dermal papillae in the plantar skin of the digits (Figure 3.38 a). Herein, $\text{Ntng}^{\text{Mech1}^+}$ peripheral endings appeared to terminate in rudimentary Meissner-form structures (Figure 3.38 b-c), which are suggested to correspond to a subset of chick Merkel corpuscles (Winkelmann and Myers, 1961; Duc et al., 1993). $\text{Ntng}^{\text{Mech1}^+}$-labeled peripheral afferents were also detected in scaly skin of the digits, but the sensory innervation could not be determined morphologically (data not shown). At the same time, $\text{Avil}^{\text{Lucy1}^+}$-driven reporter gene expression in mechanoreceptive and nociceptive peripheral projections was detected in dermal papillae and other dermal regions of the glabrous skin in digits (Figure 3.38 d-f). In analogy with previous observations, no intraepidermal termination of sensory axons was observed in chick (Figure 3.38) (Halata et al., 2003).

Altogether, STEVE-mediated stable genetic tagging of individual sensory subtypes and their circuits enables visualization and precise analysis of both central and peripheral connectivity in the chick primary somatosensory system. The peripheral innervation of $\text{Ntng}^{\text{Mech1}^+}$ Aβ-LTMRs was exclusively found in glabrous skin, possibly
terminating in rudimentary Meissner-like corpuscles, which are primarily found in featherless skin of avians. Whether Ntng1<sup>Mech<sub>1</sub></sup>-tagged peripheral termination is indeed restricted to Meissner-like corpuscles or whether Ntng1<sup>Mech<sub>1</sub></sup> projections alternatively or additionally display a topographic (‘distal-versus-proximal-limb’) innervation pattern not strictly adhering to one specific LTMR type, could not be conclusively determined. Since Herbst corpuscles are also present in glabrous skin this needs to be further addressed.

**Figure 3.38: Analysis of Ntng1<sup>Mech<sub>1</sub></sup> peripheral terminations in chick glabrous skin at E19**

(a-c) Ntng1<sup>Mech<sub>1</sub></sup>-tagged peripheral afferents terminate in dermal papillae in the plantar skin of digits in transverse E19 sections (a). Detailed view: Ntng1<sup>Mech<sub>1</sub></sup> peripheral endings innervate rudimentary Meissner-form structures (b-c). Nociceptive and mechanoreceptive axons are labeled by βIII-tubulin (TuJ1). Dotted lines demarcate epidermis and dermis. Scale bar: 25 µm.

(d-f) Avil<sup>Luy1</sup> mechanoreceptive and nociceptive peripheral afferents project in dermal papillae of glabrous skin in digits in transverse E19 sections (d). Detailed view: Avil<sup>Luy1</sup> axon endings overlap with TuJ1 (e-f). No intraepidermal termination of sensory axons is observed. Scale bar: 25 µm.
3. Results

3.8. Ntng1-NGL1 interactions in laminar targeting of sensory afferents

Precise central connectivity is a key step in the formation of well-defined neuronal circuits that relay and process sensory information. However, the transcriptional and signaling mechanisms that direct the distinct laminar target zone innervation remain poorly understood (Chen Al et al., 2006). The comprehensive analysis comprising many sensory neuron characteristics identified \( Ntng1^{Mech1} \) as a novel genetic marker, which has been shown to correspond to a discrete subclass of large-diameter \( \alpha \)-LTMRs, possibly correlating with Meissner-like LTMRs. Based on five principle observations it may be suggested that Ntng1-NGL1 interactions have a function in specific laminar targeting within the developing spinal cord of vertebrates (see Figure 1.11 b). First, \( Ntng1 \) is expressed in a subset of large-soma size somatosensory neurons in mouse DRGs (see Figure 1.11 a). Second, the selective interaction partner NGL1 appears to be expressed in the deeper laminae IV-V in mouse dorsal horn (see Figure 1.11 a). Third, the \( Ntng1^{Mech1} \) enhancer derived from the \( Ntng1 \) locus drives specific expression in a subset of \( \alpha \)-LTMRs which project to a restricted target zone within laminae IV-V in the chick dorsal horn (see Figure 3.28). Fourth, Ntng1 and NGL1 proteins have been described to be highly enriched in developing pre- and postsynaptic terminals of neurons in vertebrates (Kim et al., 2006; Brose, 2009; Woo et al., 2009). And fifth, NetrinG-NGL interactions have been implicated in excitatory synapse formation and lamina-specific segmentation of dendrites in the hippocampus (Nishimura-Akiyoshi et al., 2007; Woo et al., 2009; Seiradake et al., 2011). Consequently, the contribution of Ntng1-NGL1 interactions in laminar targeting of mechanoreceptive axons was addressed.

The impact of Ntng1-NGL1 interactions in somatosensory neuron subtypes concerning their axon projections to defined target zones in the spinal cord was examined by gain-of-function studies in late-gestation chick embryos (Chen Al et al., 2006; Guo et al., 2011). Through the combination of STEVE with the Cre/loxP system, a strategy was developed allowing specific and long-term expression of introduced transgenes at high expression levels (Figure 3.39) (Livet et al., 2007). The transgene expression cassette of the \( pCAG-loxP-ssFlag-IRES-mGFP/tdTomato-Tol2 \) is composed of the strong constitutive \( CAG \) promoter (1), a \( loxP-STOP-loxP \) cassette (2) for specific Cre
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recombinase-mediated site-specific recombination as a driver for transgene activation, a signal sequence (3) for targeting proteins to the secretion pathway, a farnesylation plasma membrane insertion signal (4), and IRES coupled reporter genes (mGFP or tdTomato) (5) ensuring bicistronic expression of an epitope-tagged transgene and the fluorescent tracer. Additionally, the transgene expression cassette is flanked by Tol2 sites (6) for transposase-mediated stable genomic integration (Kawakami and Shima, 1999; Sato et al., 2007; Takahashi et al., 2008). The mouse coding sequences of the transmembrane proteins Ntng1 and NGL1, which displayed extensive homology to the corresponding human and chick protein sequences, were introduced into the expression vector. In order to stably express NGL1 in the entire chick spinal cord, the ubiquitous Cre driver (pPGK-Cre-bpA) was used (Figure 3.39 a). Specific overexpression of Ntng1 in chick somatosensory neurons was achieved by Isl1Cre3-mediated Cre expression for selective Cre recombination in DRG neurons (Figure 3.39 b). Both transgene vectors directed intense membrane-localized transgene and reporter gene expression in all transfected neurons which was verified by detection with specific antibodies against mouse NGL1 and Ntng1 (Figure 3.40).

**Figure 3.39: Selective and stable transgene overexpression in late-gestation chick embryos**

(a) General overexpression

- **Strategy:** Subcloning of transgenes NGL1 and Ntng1 into transposon-based expression construct pCAG-loxP-ssFlag-NGL1-IRES-tdTomato-Tol2. Co-electroporation with constitutive Cre driver (PGK) and transposase (T2TP) for stable ectopic NGL1 expression in entire spinal cord (a). Co-electroporation with somatosensory-specific Cre driver (Isl1Cre3) and transposase (T2TP) for selective and stable ectopic Ntng1 expression in DRGs (b).

(b) Ntng1 overexpression

- **Strategy:** Subcloning of transgenes NGL1 and Ntng1 into transposon-based expression construct pCAG-loxP-ssFlag-Ntng1-IRES-mGFP-Tol2. Co-electroporation with constitutive Cre driver (PGK) and transposase (T2TP) for stable ectopic NGL1 expression in entire spinal cord (a). Co-electroporation with somatosensory-specific Cre driver (Isl1Cre3) and transposase (T2TP) for selective and stable ectopic Ntng1 expression in DRGs (b).
3. Results

Fig. 3.40: Verification of transgene overexpression in early stage chick embryos

(a-f) Verification of transgene expression constructs by general transgene overexpression via constitutive Cre driver (pPGK-Cre) in unilaterally transfected chick neural tube. The NGL1 expression construct shows high level expression of fluorescent protein tdTomato (a) and membrane-localized mouse NGL1 (b) in transverse neural tube sections at E6. Similarly, the Ntng1 expression construct induces robust mGFP (d) and membrane-bound mouse Ntng1 expression (e) in transfected cells of the neural tube at E6. Scale bar: 100 µm.

NGL1 is expressed in in deeper laminae of the dorsal horn presumably corresponding to laminae IV-V (Fig. 3.41 e). Hypothesizing that restricted expression of NGL1 is involved in mediating the targeting of Ntng1⁺ sensory collaterals in defined termination zones, ectopic expression of NGL1 in all laminae of the dorsal horn would be expected to alter the central connectivity pattern of Ntng1⁺ axons (Fig. 3.41 f). Therefore, NGL1 was ectopically expressed throughout all laminae of the dorsal horn and Ntng1Mech1⁻labeled central afferent collaterals were analyzed for their laminar targeting. Robust expression of NGL1 and fluorescent reporter protein in all laminae of the dorsal horn was achieved (Fig. 3.41 d), whereas dorsal horn laminar organization appeared unaffected (Fig. 3.41 c). Like in control experiments, Ntng1Mech1⁺ projections penetrated the superficial dorsal horn at the expected medial position and terminated in the ventromedial quadrant presumably corresponding to laminae IV-V.
3. Results

(compare Figure 3.41 a-c and Figure 3.28 c-d). No obvious shift in the distribution of Ntng1Mech1+ central collaterals to other NGL1 expressing laminae was detected.

Alternatively, if Ntng1-NGL1 interaction indeed was a determinant for the specific termination pattern in LTMR circuits, the status of Ntng1 expression in sensory neurons should influence the positioning of their collaterals within the spinal cord (Figure 3.42 f). Ectopic expression of Ntng1 in prospective nociceptive and proprioceptive sensory neurons could therefore alter their axonal projections to adopt Ntng1+ LTMR-like features and to prompt axons to assume a medioventral position within the dorsal horn (laminae IV-V), which defines a region where interaction partner NGL1 is expected to be accumulated (Figure 3.42 g). Therefore, Ntng1 was overexpressed in all somatosensory neuron subtypes and the central innervation of...
transfected neurons was observed. In congruence with the NGL1 overexpression analysis, no drastic influence of forced Ntng1 expression in the axonal trajectories of nociceptive, mechanoreceptive or proprioceptive neurons was observed (compare Figure 3.42 a-e and Figure 3.15 a-e).

In summary, Ntng1 Mech1+ central collaterals were not redirected into different NGL1 expressing laminae by ectopic NGL1 expression throughout the dorsal horn.
(Figure 3.41 g). In addition, ectopic expression of Ntng1 in all subtypes of somatosensory neurons did not change the subtype-specific laminar targeting pattern of nociceptive, mechanoreceptive and proprioceptive neurons (Figure 3.42 h). Hence, it can be concluded that, at least under conditions of forced expression in chick, neither Ntng1 nor its interaction partner NGL1 is sufficient to alter the overall central connectivity of axonal projections in the spinal cord. However, it cannot be excluded that Ntng1-NGL1 interactions play a role in fine-grained changes in laminar and sublaminar targeting, like in the lamina-specific differentiation of dendrites. Through a novel in silico-to-in vivo screen for neuron subtype-specific gene regulatory elements, a unique LTMR circuit defined by Ntng1 activity was uncovered. This circuit appears to relay touch responses from Meissner-like glabrous skin Aβ-LTMRs to laminae IV-V in the dorsal horn. However, deciphering the precise roles of endogenous Ntng1 activity in LTMRs, and that of its putative interaction partner NGL1 in the dorsal horn, during somatosensory circuit assembly or function remains to be addressed.
4. Discussion

4.1. STEVE-mediated stable genetic tagging of somatosensory neurons

The sense of touch is essential for a variety of behaviors in virtually all higher animal species (Lumpkin et al., 2010). Formation and specification of precise sensory neuronal circuits during embryogenesis represents a crucial step in the development of a functional nervous system (Kandel et al., 2000). However, little is known about molecular mechanisms underlying sensory neuron specification, the unique function of specialized sensory neurons, and the logic of sensory circuit organization, which is mainly due to the limited availability of sensory neuron subtype-specific markers facilitating their comprehensive analysis (Marmigère and Ernfors, 2007; Ma, 2009; Lumpkin et al., 2010; Nilius, 2010; Li et al., 2011; Lallemend and Ernfors, 2012). This study aimed at the identification of novel genetic markers for distinct sensory subtypes to provide more detailed insights into sensory neuron circuit organization and developmental assembly. These overall aims were addressed by screening for sensory neuron subtype-specific enhancer activities in a novel chick model system.

An extensive enhancer analysis in rodent transgenic animals, including screening of multiple genomic sequences for enhancer activities in vivo, implicates high costs, resource-intensiveness and lengthy generation times (Timmer et al., 2001). The chicken as a complementary model system shares developmental and structural similarities to mammals, including universal conserved gene regulatory mechanisms, and provides a straightforward and rapid in vivo transfection with minimal time effort and low costs (Novitch et al., 2001; Thaler et al., 2002; Lee SK et al., 2004; Dasen et al., 2005; Uchikawa, 2008). Therefore, in ovo electroporation greatly facilitated an enhancer activity screen along with the study of sensory neuron specification and connectivity in vertebrates. A straightforward strategy was explored based on vertebrate transposon-based transgenesis, allowing for the first time effective, selective and stable genetic tagging of somatosensory neurons and their connectivity patterns in late-gestation chick embryos. The designed stable expression vector STEVE, including the minimal TATA box promoter and fluorescent reporter genes, provided the basis for a sensitive enhancer analysis by facilitating ready detection of
putative enhancer activities in vivo. Persistent expression in transfected cells at later developmental stages along with a long-term observation of enhancer activities was ensured by Tol2 transposon-based random genomic integration. This was particularly necessitated by the high mitotic rates of neural crest cells (NCCs), causing non-integrated transgenes to disappear rapidly by dilution and progressive degradation (Kawakami and Shima, 1999; Sato et al., 2007; Takahashi et al., 2008; Yokota et al., 2011). For high transfection efficiency in the sensory lineage, in ovo electroporation of the chick neural tube was consequently performed between Hamburger-Hamilton (HH) stages 11 and 13 before NCC emigration (Hamburger and Hamilton, 1951; Serbedzija et al., 1990; Kasemeier-Kulesa et al., 2005; Krispin et al., 2010). Previous studies reported of biased transfection efficiency within the somatosensory lineage towards a particular somatosensory subtype (Chen AI et al., 2006; George et al., 2007). In contrast, STEVE provided unbiased stable transfection and expression in the entire spinal cord and dorsal root ganglia (DRGs), as verified with the constitutive CAG promoter, whereas the enhancer-less construct did not display activity in neural tissue. The effective method of in ovo electroporation for introducing exogenous DNA in combination with STEVE therefore created the prerequisites for the detection of cis-regulatory activities and their subsequent use for the stable genetic tagging of somatosensory neurons in late-gestation chick embryos that were inaccessible to previous methods.

4.2. Establishment of Isl1Crest3 and AvilLucy1 as pan-sensory neuron markers

In order to identify novel pan-sensory and sensory subtype-specific markers literature and gene expression data bases were screened for putative candidates expressed by DRG neurons (Yin et al., 2002; Uemura et al., 2005; Alvarez-Bolado and Eichele, 2006). Genomic comparison of chosen candidate gene loci between multiple species revealed a wide range of evolutionary conserved non-coding regions (ECRs) (Ovcharenko et al., 2004). Based on homology and conservation of putative transcription factor binding sites the most promising ECRs were analyzed. It was found that five out of seven tested ECRs were driving reporter gene expression in neuronal
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tissue of the chick, with both specific and non-specific activities (see Table 3.1). The general sensory neuron-specific enhancers \( \text{Isl1}^{\text{Crest3}} \) and \( \text{Avil}^{\text{Lucy1}} \), as well as the mechanoreceptive neuron-specific enhancer \( \text{Ntng1}^{\text{Mech1}} \) were discovered via this method by recapitulating the endogenous expression pattern of the corresponding genes in DRG neurons. These results confirmed the efficacy of the designed screening strategy for the rapid \textit{in silico-to-in vivo} enhancer identification (Boffelli \textit{et al.}, 2004; Uchikawa, 2008; Visel \textit{et al.}, 2008). Additionally, the method enabled a significantly easier access to sensory neuron manipulation studies in both early and late-gestation embryos, compared to previous manipulation approaches (Chen \textit{et al.}, 2006; George \textit{et al.}, 2007). The analysis of identified ECRs supported previous observations that enhancers can modulate gene expression independently of their orientation or their distance to the gene transcription start (Khoury and Gruss, 1983; Blackwood and Kadonaga, 1998; Pennacchio \textit{et al.}, 2006; Doh \textit{et al.}, 2007). \( \text{Isl1}^{\text{Crest3}} \), for example, originally located more than 323 kb upstream of the gene transcription start, was subcloned just upstream to the reporter gene coding sequence in \textsc{steve} with any functional impairment, thereby demonstrating the relative independence between enhancer distance and its function in gene regulation. In accordance with previous studies, a high ECR sequence conservation is reflected by a high probability of enhancer activity of mouse genome-derived ECRs in the chick embryo, thus emphasizing the idea that mammals and avians share many basic gene regulatory mechanisms in the primary somatosensory system (Timmer \textit{et al.}, 2001; Uchikawa \textit{et al.}, 2004; Dasen \textit{et al.}, 2005).

In this study, \( \text{Isl1}^{\text{Crest3}} \) and \( \text{Avil}^{\text{Lucy1}} \) were established as new pan-sensory neuron markers by driving a specific and strong reporter gene expression in somatosensory neurons from embryonic stages into adulthood in chick. Along with this, the pan-sensory markers provided both central and peripheral tagging of sensory axon collaterals and provided the feasibility to directly analyze the connectivity of somatosensory neurons throughout late gestation in chick embryos. The overall developmental sequence of nociceptive, mechanoreceptive and proprioceptive collateral extension into the gray matter in chick mirrored that observed in the mammalian spinal cord. However, the delay in the establishment of nociceptive compared to proprioceptive afferent connectivity observed in mammals appeared
temporally compressed in avians, likely reflecting an ontogenic adaptation in precocial birds (Fitzgerald, 1987; Mirnics and Koerber, 1995; Ozaki and Snider, 1997; Marmigère and Ernfors, 2007). Furthermore, these observations provided evidence that central connectivity in the spinal cord is established by a majority of central afferent collaterals that directly home in on their respective target laminae (Eide and Glover, 1997; Fitzgerald, 1987; Davis et al., 1989; Ozaki and Snider, 1997; Redmond et al., 1997; Fitzgerald, 2005).

As an exception to the above mentioned trend, the examined ECRs derived from the well-established nociceptive marker TrkA and the proprioceptive marker Runx3 did not reflect the endogenous gene expression in sensory DRG neurons, and showed no or unspecific enhancer activity in the central and peripheral nervous system. Since many genes are regulated by complex arrays of enhancers for temporal and spatial gene expression, the discovery of the particular enhancers being responsible for sensory neuron-specific gene expression may not always be straightforward, often requiring more extensive experimental trials (Pennacchio et al., 2006; Alberts et al., 2008). Although highly conserved, no enhancer activity of TrkAECR1 and Runx3ECR3 could be observed in neuronal tissue of the embryonic chick. One might speculate that these sequences might still include functional regulatory elements which could mediate enhancer activity in different tissue not accessible via chick neural tube electroporation or become active at stages that were not analyzed (Uchikawa et al., 2003; Uchikawa, 2008; Visel et al., 2009). Silencer and insulator elements in a gene locus indispensable for mediating a specific gene expression are, like enhancer elements, thought to be evolutionary highly conserved. Since it is impossible to distinguish between enhancer, silencer and insulator elements in a screen for ECRs, it could be possible that the chosen ECRs are silencers or insulators, which are not sufficient to drive a gene transcription (Blackwood and Kadonaga, 1998; Woolfe et al., 2005; Visel et al., 2009). The Runx3ECR1- and Runx3ECR2-driven non-specific expression in different subtypes of sensory DRG neurons in the late-gestation chick was not reflecting the endogenous gene expression normally restricted to the proprioceptive subtype. It is possible that the ECR-mediated expression becomes confined to the proprioceptive neuron subtype with additional enhancer and silencer elements which are necessary in the genomic locus to specify Runx3 expression (Lee SK et al., 2004; Doh et al.,
By analysis of further candidates, additional enhancers mediating expression in different somatosensory subtypes could be discovered. A comprehensive analysis with a variety of diverse sensory subtype-specific markers could provide a better understanding how specification, as well as precise circuitries of the different sensory neuron classes are established (Lallemend and Ernfors, 2012).

The ability to directly relate sensory identities in chick to specific sensory modalities in other vertebrate species is currently limited by the comparatively poorly resolved structure-function relationships in the avian somatosensory system. The improvement of this situation is, however, expected to go hand-in-hand with the ongoing identification and characterization of molecularly defined sensory neuron subclasses. By allowing systematic exploration of molecularly defined neuronal identities, the STEVE-based screening for cis-regulatory activities in chick offers a potentially powerful pre-screening platform for targeting discrete sensory neuronal populations in other vertebrate systems by uncovering principles of their specification and connectivity. While in many cases the mouse will likely remain the gold standard for the genetic dissection of neuronal circuits, the strategic use of auxiliary models facilitating rapid molecular identification and interrogation of neuronal identities could thereby accelerate the drive to map mammalian connectomes.

4.3. Neurogenic equivalence of ipsilaterally and contralaterally migrating neural crest cells

During early embryonic development, NCCs emigrate from the neural tube towards their respective peripheral targets, whereby some NCCs cross the neural tube midline and contribute to the contralateral NCC migratory stream (Serbedzija et al., 1990; Frank and Sanes, 1991; Carmona-Fontaine et al., 2008; Squire et al., 2008). George and colleagues reported of lineage-restricted premigratory NCCs in diametrically opposed (ipsilateral and contralateral) migratory streams, such that contralaterally migrating NCCs are biased to generate nociceptive somatosensory neurons (George et al., 2007; Lefcort and George, 2007; George et al., 2010). The potential presence of
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lineage-restricted NCC populations with unique migratory trajectories has far-reaching implications for understanding NCC biology, including the origin of pain-sensing circuits, and could have profound consequences for experimental strategies targeting ipsilateral or contralateral NCC derivatives.

Through STEVE-mediated direct cell lineage-tracking and systematic analysis of neuronal NCC progeny and their circuitries in chick conclusive evidence could be provided that ipsilateral and contralateral NCC streams are neurogenically equivalent and give rise to the same principal set of NCC derivatives, including peripheral neuron types and their circuitries. It is likely that the use of stable somatic transgenesis for tracking NCC lineages effectively sidestepped past limitations inherent to conventional plasmid-based transfection, thus, contributing to the different outcomes between this study and preceding experiments (George et al., 2007). The limited intracellular half-life of non-integrating plasmid DNA confines lineage-tracking experiments to largely immature NCC progeny in DRGs (Yokota et al., 2011). Moreover, in analogy to classical dye-tracing procedures, this might collude with fluctuating tracing efficacies to skew outcomes towards certain cell classes. This could be caused by disparate dilution rates in sub-lineages progressing through few or several cell divisions, especially in the highly proliferative NCC lineages (Stern and Fraser, 2001; Lacar et al., 2010). These effects would have previously eluded detection by the omission of a direct comparative analysis of ipsilaterally- or contralaterally-derived DRG neuron classes, which is a prerequisite for conclusively assessing any neurogenic bias (or lack thereof) of ipsilateral or contralateral NCCs.

Based on obtained observations of STEVE-mediated unbiased bilateral targeting of DRG neurons, it is proposed that the entry of a contralateral migratory trajectory by some NCCs is most parsimoniously explained by stochastic events, likely resulting from homotypic interactions driving the initial pattern of NCC emigration (Carmona-Fontaine et al., 2008). Both, ipsilateral and contralateral migratory streams would thereby derive from a common pool of precursors subjected to the same spatiotemporal factors influencing NCC potential (Le Douarin and Kalcheim, 1999; Anderson, 2000; Harris and Erickson, 2007; Krispin et al., 2010). A practical consequence of this is the emerging possibility of using unbiased targeting of
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contralateral NCC progeny, such as for the sparse labeling of peripheral neurons for in-depth circuit mapping (Jefferys and Livet, 2012; Chen Al et al., 2006). This feature was utilized for the analysis of somatosensory subtype-specific central trajectories in the developing chick embryo (see Figure 3.23-3.26).

4.4. A novel subclass of touch receptor neurons defined by $Ntng1^{Mech1}$

Dependent on the specific functional characteristics associated with the detection of different touch stimuli, low-threshold mechanoreceptors (LTMRs) can be broadly subdivided in heavily myelinated rapidly adapting (RA) $\alpha$-LTMRs, slowly adapting (SA) $\alpha$-LTMRs, lightly myelinated $\alpha$-LTMRs and unmyelinated C-LTMRs. The medium-diameter $\alpha$-LTMRs and the small-diameter C-LTMRs differ morphologically and physiologically from the large-diameter $\alpha$-LTMRs of both RA and SA types (Lewin and Moshourab, 2004; Lumpkin and Caterina, 2007; Ma, 2009). By demonstrating the feasibility of the STEVE system for the rapid identification and interrogation of neuronal subtype identities, $Ntng1^{Mech1}$ was shown to label a novel molecularly defined subset of sensory neurons whose anatomical, morphological, electrophysiological, functional and molecular characteristics correspond to a discrete subclass of $\alpha$-LTMRs. Based on the similar $Ntng1^{Mech1}$-mediated expression pattern in large-size sensory neurons and its close genomic localization to the gene transcription start, $Ntng1^{Mech1}$ is expected to correspond to an enhancer associated with the $Ntng1$ gene and consistently to reflect the endogenous gene expression in sensory DRG neurons (Yin et al., 2002; www.brain-map.org). Nevertheless, a final validation for a direct association should be ultimately conducted via two-color fluorescence in situ hybridization analysis against the $Ntng1^{Mech1}$-driven reporter gene and endogenous Ntng1 or via co-expression analysis with specific antibodies against the reporter gene and chick Ntng1 (Levsky and Singer, 2003).

In order to classify $Ntng1^{Mech1+}$ DRG neurons according to defined sensory neuron subtypes a comprehensive analysis addressing typical characteristics of sensory neuron subclasses was performed. $Ntng1^{Mech1}$ consistently labeled a population of
large-diameter sensory neurons clustered in the ventrolateral position of the DRGs, which is mainly occupied by mechanoreceptive and proprioceptive neurons in chick (Frank and Sanes, 1991; Harris and Erickson, 2007). More strikingly, Ntng1\textsuperscript{Mech1+} central axon collaterals projected into a highly confined region in the medial dorsal horn of the spinal cord which is the target zone of LTMRs in chick (Eide and Glover, 1997; Wild et al., 2010). Furthermore, Ntng1\textsuperscript{Mech1+}-labeled sensory collaterals separated from nociceptive central afferents targeting laminae I-II in the lateral dorsal horn and from proprioceptive afferents projecting to the intermediate and ventral spinal cord (Scott, 1992; Chen Al et al., 2006; Lallemend and Ernfors, 2012). By labeling the largest DRG neurons which are highly myelinated based on the co-expression with NF200, the properties of Ntng1\textsuperscript{Mech1+} neurons highly correlated with \(\text{A}\beta\)-LTMR characteristics (Harper and Lawson, 1985; Lechner et al., 2009). Based on these criteria, Ntng1\textsuperscript{Mech1+}-tagged sensory neurons strongly differed from small-size lightly myelinated or unmyelinated nociceptive neurons, as well as from medium-diameter myelinated \(\text{A}\delta\)-LTMRs and small-diameter unmyelinated C-LTMRs (Harper and Lawson, 1985; Marmigère and Ernfors, 2007). The electrophysiological analysis revealed that Ntng1\textsuperscript{Mech1+} DRG neurons represent a discrete population of sensory neurons with spike characteristics and mechanosensitive currents typical of embryonic LTMRs, concomitant with the exclusion of functional properties characteristic for nociceptive neurons (Koerber et al., 1988; Djouhri et al., 1998; Fang et al., 2005; Lechner et al., 2009). The stereotypic laminar connectivity pattern established by Ntng1\textsuperscript{Mech1+} DRG neurons displayed a selective termination within a medioventral domain presumably corresponding to laminae IV-V by excluding lamina III of the dorsal horn. Recent studies showed that the central collaterals of molecularly identified \(\text{A}\beta\)-LTMRs, including both RA and SA types, broadly target laminae III to V, which is consistent with the connectivity patterns of morphologically or physiologically identified LTMRs in both, mammals and birds (Necker, 1990; Eide and Glover, 1997; Bourane et al., 2009; Luo et al., 2009; Honma et al., 2010; Li et al., 2011; Lallemend and Ernfors, 2012). Moreover, transganglionic tracing of morphologically identified skin mechanoreceptors in rat have previously revealed that \(\text{A}\delta\)-LTMR afferents tend to be concentrated in laminae II (inner) and III (Light and Perl, 1979; Li et al., 2011). It may therefore be tempting to speculate that \(\text{A}\delta\)-LTMRs are Ntng1\textsuperscript{Mech1-}. The exclusion of
laminae I-III and restriction to laminae IV-V by Ntng1 Mech1+ collaterals thus appears to uncover a novel molecularly defined subclass of Aβ-LTMRs (Figure 4.1).

In analogy to the fine-grained laminar and sublaminar segregation patterns of molecularly defined nociceptor identities, the functional specialization of LTMRs could thus in general be expressed as unique labeled lines relaying submodalities to discrete laminal targets (Zylka et al., 2005; Li et al., 2011). Currently, it is generally believed that LTMRs arise from two early populations of mechanoreceptive progenitors expressing either Ret (earlyRet+ population) or TrkB (TrkB+ population) (Lallemend and Ernfors, 2012). The earlyRet+ lineage further subdivides which finally results in five molecularly unique lineages: Ret+/MafA+ (1), Ret+/MafA+/TrkB+ (2), Ret+/MafA+/TrkC+ (3), Ret+/TH+ (4) and TrkB+ (5) (Bourane et al., 2009; Luo et al., 2009; Honma et al., 2010; Abdo et al., 2011; Heidenreich et al., 2011; Li et al., 2011; Wende et al., 2012). In mammals, it has been speculated that the Ret+/MafA+ subsets (lineages 1-3) of LTMR neurons could predominantly represent RA Aβ-LTMRs, the Ret+/TH+ subset (lineage 4) is predicted to label C-LTMRs, whereas the TrkB+ population (lineage 5) was associated with Aδ-LTMRs (Bourane et al., 2009; Luo et al., 2009; Heidenreich et al., 2011; Li et al., 2011; Wende et al., 2012). Based on
current literature no specific markers for SA Aβ-LTMRs have been identified yet. However, in most cases the correlation between the molecularly unique LTMR subtypes with distinct mechanoreceptive end organs needs to be determined in more detail. Additionally, the regulatory mechanisms regarding specification and connectivity of the different LTMR subtypes still remain unknown (Lallemand and Ernfors, 2012). Since the early development of sensory differentiation displays homology in vertebrates, it might be suggested that also later specification and maturation of LTMRs is highly conserved between mammals and avians (Koltzenburg and Lewin; 1997; Marmigère and Ernfors, 2007; Guo et al., 2011).

In the present study, it was found that the Ntng1\textsuperscript{Mech1}\textsuperscript{-}-tagged DRG neuron population partially overlaps with earlyRet\textsuperscript{*} (lineages 1-4) and TrkB\textsuperscript{*} (lineages 2+5) mechanoreceptive neurons, with 31% of Ntng1\textsuperscript{Mech1+} neurons co-expressing Ret and 33% co-expressing TrkB. By assumption of no co-expression between Ret and TrkB in any of the LTMR lineages, which is actually present in lineage 2, at least 36% of Ntng1\textsuperscript{Mech1+} neurons express neither Ret nor TrkB. This indication therefore points towards a molecularly novel Ret\textsuperscript{-} TrkB\textsuperscript{-} LTMR subtype defined by Ntng1\textsuperscript{Mech1} (additionally to lineages 1-5), which has not yet been described. Based on this, it would be predicted that Ntng1\textsuperscript{Mech1+} Aβ-LTMRs are partially independent of the Neurotrophin Ret/Trk receptor signaling during development, which may be further clarified with a DRG explant culture assay (Bilsland et al., 1999; Bourane et al., 2009; Lechner et al., 2009; Wang and Marquardt, 2012). Accordingly, by incubation of Ntng1\textsuperscript{Mech1}-transfected sensory neurons with single growth factors (GDNF, NGF, BDNF, NT-3) the influence on differentiation, neurite outgrowth and survival of Ntng1\textsuperscript{Mech1+} neurons could be analyzed in order to classify the Ntng1\textsuperscript{Mech1} population more precisely (Figure 4.2). Analogous to the division of earlyRet\textsuperscript{* } mechanoreceptors into diverse subclasses, Ntng1\textsuperscript{Mech1+} neurons may thus be tentatively divided into Ntng1\textsuperscript{Mech1+} (1’), Ntng1\textsuperscript{Mech1+}/Ret\textsuperscript{*} (2’), Ntng1\textsuperscript{Mech1+}/TrkB\textsuperscript{*} (3’) and Ntng1\textsuperscript{Mech1+}/TrkC\textsuperscript{*} (4’) Aβ-LTMRs, whereas a potential overlap is considered to be possible between the Ret\textsuperscript{*} and TrkB\textsuperscript{*} subsets (2’+3’), as well as between the Ret\textsuperscript{*} and TrkC\textsuperscript{*} populations (2’+4’). However, given the apparent molecular heterogeneity of the larger class of Aβ-LTMRs, and the smaller class of Ntng1\textsuperscript{Mech1+} ‘laminae IV-V’ Aβ-LTMRs, it is also
possible that the relationship between mechanosensory neuron subclass and Ret/Trk receptor expression is even more complex.

In mammals, the peripheral collaterals of RA Aβ-LTMRs terminate in hair follicles, as well as Pacinian and Meissner corpuscles that dynamically respond to light touch, vibration, flutter and skin movements, while SA Aβ-LTMRs innervate Merkel discs and Ruffini corpuscles that relay static responses to dermal stretch and indentation (Lewin and Moshourab, 2004; Lumpkin and Caterina, 2007; Ma, 2009). The Aδ-fiber D-hair receptors and C-fiber LTMRs are associated with hair follicles (Lumpkin et al., 2010; Li et al., 2011). In contrast to the well characterized sensory end organs in mammals there is only limited information on avian mechanoperception and its adaptations in feathery versus hairy skin. In birds, detection of different touch stimuli appears to be mainly mediated by Herbst and Merkel corpuscles (Gottscheidt 1985; Necker 2000). Herbst corpuscles are considered as the avian equivalent of the mammalian Pacinian corpuscles, which respond to vibration and acceleration (Gottscheidt, 1985; Necker, 2000). The corpuscles are widely distributed throughout the feathery skin by forming terminals in the collar region of most feather follicles, but also innervate the dermis of glabrous skin (Duc et al., 1993). Merkel cell receptors are primarily found in featherless skin of avians, where they respond to sustained pressure (Necker, 2000; Halata et al., 2003). In contrast to mammals, Merkel cells here differ from mammalian
Merkel cell-neurite complexes by accumulating and forming rudimentary Meissner-like corpuscles exclusively present in the superficial layer of the dermis and being highly enriched in dermal papillae (Winkelmann and Myers, 1961; Duc et al., 1993). The electrophysiological classification of Herbst and Merkel corpuscles among defined RA or SA modalities is complicated by conflicting findings, with some groups reporting that both receptor types are RA, whereas others identified both RA and SA responses (Necker, 1985; Gentle, 1989; Necker, 1990; Duc et al., 1993). The peripheral innervation of Ntng1 Mech1+ Aβ-LTMRs was exclusively found in plantar and scaly regions of the glabrous skin. The innervation of dermal papillae to rudimentary Meissner-like structures indicates that Ntng1 Mech1+ axons terminate in this subset of avian Merkel corpuscles. Whether the Ntng1 Mech1+ and Ntng1 Mech1- neurons are indeed tuned to specific mechanosensory submodalities awaits further study, however.

A co-labeling with sensory end organ specific markers could provide more precise conclusions of the Ntng1 Mech1+-mediated target specificity (Duc et al., 1993). Since Herbst corpuscles are also present in glabrous skin it need to be determined whether the peripheral termination of Ntng1 Mech1-tagged LTMRs is restricted to Merkel corpuscles or whether these projections alternatively or additionally display a topographic ‘distal-versus-proximal-limb’ innervation pattern. However, such a ‘distal-versus-proximal-limb’ regional specificity of the Ntng1 Mech1 peripheral projections appears unlikely, as a topographic skin innervation pattern is generally thought to be reflected by central afferents that organize in column-like structures from medial to lateral throughout different laminae of the dorsal horn (Wild et al., 2010; Li et al., 2011). Since Ntng1 Mech1+ central afferent collaterals occupy the entire medioventral domain of the dorsal horn presumably corresponding to laminae IV-V by excluding laminae I-III, Ntng1 Mech1 is rather more likely restricted to a specific LTMR subtype. Due to the so far limited morphological and electrophysiological characterization of LTMR subtypes in avians and due to the difficulty of pinpointing the homologous mammalian sensory end organs, conclusive statements regarding the stereotypic sensory end organ innervation of Ntng1 Mech1+ LTMRs remain challenging. In order to address the issue of a precise electrophysiological characterization of the avian LTMR subtypes, including Ntng1 Mech1-tagged neurons, particularly with regard to their response properties to different stimuli, in vitro skin nerve preparations will have to be
performed (Lewin and Moshourab, 2004; Heidenreich et al., 2011; Wende et al., 2012). Therefore, a novel not-yet-described strategy was designed to selectively tag and record single axons of a defined population in the chick. Since the avian transient receptor potential cation channel V1 (TrpV1), in contrast to the mammalian form, is insensitive to capsaicin, mammalian TrpV1 will be selectively expressed in Ntng1\textsuperscript{-Mech1+} peripheral axons, which then can be recognized after capsaicin exposure of the skin (Wood et al., 1988; Jordt and Julius, 2002). Subsequently, the responses of Ntng1\textsuperscript{-Mech1+} axons can be recorded after variable mechanical stimulation of the skin (vibration, sustained pressure, etc.) in order to classify the LTMR population among defined RA or SA modalities. Isl1\textsuperscript{Cre+/-} or Avil\textsuperscript{Luc+/-}-driven TrpV1 in all somatosensory subtypes might serve as a reference for the different response properties present in the chick somatosensory circuitry.

Finally, the overall aim for the near future is to recapitulate, verify and to in depth characterize the Ntng1\textsuperscript{-Mech1+}-mediated LTMR subtype-specific expression in the mouse via specific antibodies against Ntng1, as well as by stable transgenesis, as the mouse serves as a well-established model system in regard to study the development and the function of somatosensory circuitries (Luo et al., 2009; Heidenreich et al., 2011). While there is a lack of information regarding the specific mechanosensory modality they relay, Ntng1\textsuperscript{-Mech1+} neurons constitute a novel molecularly unique subclass of A\(\beta\)-LTMRs. Moreover, the accelerated identification of molecularly defined somatosensory neuron subclasses by STEVE-based transgenesis in chick may accelerate the comprehensive mapping of neuronal identities associated with mechanosensory (sub)modalities.

4.5. NetrinG-NGL interactions in mechanoreceptive circuit formation

Ntng1\textsuperscript{-Mech1} defines a novel subclass of large-diameter A\(\beta\)-LTMRs whose central collaterals selectively terminate within a defined domain of the dorsal horn presumably corresponding to laminae IV-V. Since NetrinG-NGL interactions have been implicated in excitatory synapse formation and lamina-specific segmentation of dendrites in the
Ntng1 expression is expected to be restricted to the deeper laminae of the dorsal horn presumably corresponding to laminae IV-V. If Ntng1-NGL1 interactions would be the determinants for laminar targeting of Ntng1+ sensory collaterals in their defined termination zone of the dorsal horn, ectopic expression of NGL1 in all laminae should be able to alter the central connectivity pattern of Ntng1+ axons. Alternatively, ectopic expression of Ntng1 in prospective nociceptive and proprioceptive sensory neurons should be able to alter their axonal projections to adopt Ntng1+ LTMR-like features and to redirect their axons towards regions in the dorsal horn where interaction partner NGL1 is expected to be accumulated. Primary experiments showed that Ntng1Mech1+ central collaterals were not redirected into different NGL1 expressing laminae by ectopic NGL1 expression throughout the dorsal horn. Furthermore, no change in the phenotypic laminar targeting of Ntng1 expressing nociceptive, mechanoreceptive and proprioceptive neurons was observed. These primary observations lead to the conclusion that Ntng1-NGL1 interactions are not sufficient to alter the stereotypic laminar targeting of somatosensory projections in the spinal cord. However, it cannot be excluded that Ntng1-NGL1 interactions might be involved in fine-grained changes in laminar and sublaminar connectivity, in analogy to the lamina-specific dendrite segmentation in hippocampal neurons observed by previous studies (Nishimura-Akiyoshi et al., 2007; Seiradake et al., 2011). The design of a more precise and more sensitive analysis, which facilitates the detection of changes at sublaminar levels, is essential to reach the final conclusion regarding the Ntng1-NGL1 function in central targeting of LTMR afferents. In vitro studies might address the individual influence of Ntng1-NGL1 interactions in axon guidance of Aβ-LTMR neurons with less interference factors, compared to in vivo studies. The stripe assay, for example, has been proven to be powerful in uncovering guidance cues and mechanisms in many principal neuronal projections (Knöll et al., 2007; Kao and Kania, 2011). Consequently,
4. Discussion

Ntng1Mech1-tagged somatosensory neurons in DRG explant cultures could be analyzed for their extending neurites along ligands printed in a striped pattern in order to study the responses of growing axons to Ntng1-NGL1 signaling. Furthermore, coculture of Ntng1Mech1-transfected DRG explants with NGL1 expressing cell aggregates could address the attractive or repulsive Ntng1-NGL1 interaction present in LTMR afferents (He and Tessier-Lavigne, 1997; Anderson et al., 2003). Ntng1-NGL1 adhesion might also be involved in peripheral targeting of Aβ-LTMR collaterals. Since the analysis of peripheral skin innervation in avians is difficult, due to limited information about the mechanoreceptive end organ organization, transgenic mice, carrying the Ntng1Mech1 expression cassette, could bridge the gap between the molecular identity of the Aβ-LTMR subpopulation and their precise peripheral and central circuit formation.

The assembly of neuronal circuits relies on a complex interplay of diverse combinations of different adhesion systems that work in parallel and trigger the precise cell-type specific connectivity (Brose, 2009; Lallemend and Ernfors, 2012). Dysfunction of one component, for example Ntng1-NGL1 interactions in LTMR circuitries, may therefore not necessarily result in a drastically altered connectivity, because of remaining and partially redundant factors. Interestingly, the NetrinG family member Ntng2 and its ligand NGL2 display a comparable expression pattern like Ntng1 in large-diameter DRG neurons and NGL1 in the deeper laminae of the dorsal horn, respectively (Yin et al., 2002; www.brain-map.org). If both Ntng1 and Ntng2 are co-expressed in the same LTMR population, these highly homologous factors would be likely to act in a redundant manner to achieve laminar targeting specificity. Another possibility might be, in analogy to their roles in the organization of lamina/pathway-specific dendrite differentiation in the hippocampal neurons, that Ntng1 and Ntng2 are expressed in different somatosensory subpopulations mediating specific central innervation in discrete domains of the spinal cord. The study of Ntng1/Ntng2 double knockout mice could reveal new insights regarding these issues (Nishimura-Akiyoshi et al., 2007). In the future, it will be interesting to decipher more precisely, which additional factors provoke the differentiation towards the Ntng1Mech1-specific subclass of large-diameter Aβ-LTMRs, how regulatory mechanisms mediate the unique organization of this circuit and what kind of touch responses are relayed by the Ntng1Mech1+ Meissner-like glabrous skin LTMRs to laminae IV-V in the dorsal horn.
5. Summary

The assembly of circuitries for somatosensory perception relies on a complex interplay of neuronal diversification, specification, phenotypic modulation and establishment of precise connectivity patterns during embryogenesis. However, little is known about the molecular mechanisms controlling the establishment of well-defined somatosensory connectivity, which is to a large part due to the limitation of sensory neuron subtype-specific molecular markers. This study aimed at the identification of novel genetic markers for distinct somatosensory neuron subtypes to provide more detailed insights into primary somatosensory circuit formation, organization and function. These aims were addressed by a newly established in silico-to-in vivo screen for neuron subtype-specific enhancer activities in the chick.

The designed strategy, based on the stable expression system STEVE, facilitated the detection of cis-regulatory activities and their subsequent use for the rapid, efficient and stable genetic tagging of somatosensory neurons in chick embryos at late-gestation stages that were inaccessible to previous methods. The feasibility of the system for the rapid identification and interrogation of sensory neuronal identities was demonstrated by the establishment of Isl1Crest3 and AvilLucy1 as pan-somatosensory neuron markers. These newly identified enhancers mediated a specific and robust reporter gene expression in somatosensory neurons from embryonic stages at least until hatching and facilitated both central and peripheral visualization of sensory axon collaterals. A detailed reconstruction of the sensory trajectories regarding their phenotypic laminar targeting revealed new insights in the complex central connectivity of the different somatosensory lineages. Furthermore, STEVE-based direct lineage-tracking and systematic analysis of neural crest cell progeny in chick provided conclusive evidence that ipsilateral and contralateral neural crest streams are neurogenically equivalent, giving rise to the same principal set of derivatives, including peripheral neuron types and their circuitries.

Through the selective screen for sensory neuron subtype-specific gene regulatory elements and the subsequent comprehensive analysis comprising many sensory neuron characteristics, Ntng1Mech1 was identified as a novel genetic marker for a
discrete subclass of large-diameter $A\beta$ low-threshold mechanoreceptors. Ntng1 activity thereby uncovered a novel circuit that appears to relay touch responses from Meissner-like glabrous skin mechanoreceptive neurons exclusively to laminae IV-V in the dorsal horn. Ongoing work is directed at deciphering the precise roles of endogenous Ntng1 activity in this subclass of $A\beta$ low-threshold mechanoreceptors, and that of its putative interaction partner NGL1 in the dorsal horn, during somatosensory circuit assembly.

Taken together, this study designed a simple screening strategy that permits rapid, efficient and stable genetic tagging of neuronal subtypes \textit{in vivo}, which thus provides the basis for the systematic discovery of late-onset sensory neuron subtype-specific gene regulatory activities. Thereby, a novel molecularly defined subclass of touch receptor neurons was uncovered, thus providing the utility of the system for uncovering the neural basis of discrete sensory modalities relayed by the primary somatosensory system. In addition, the system facilitated stable cell fate-tracking, which unraveled outstanding features of the primary somatosensory neuron lineage. Besides facilitating the genetic dissection of the somatosensory system, the strategy also offers an effective pre-screening platform for targeting genetically identified neuronal subtypes in other vertebrate species.
6. References


6. References


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8. Curriculum Vitae

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9. Declaration

I hereby declare that I prepared this thesis, entitled “Exploring the neural basis of touch through selective and stable genetic tagging in the chick somatosensory system”, on my own and with no other sources and aids than quoted.

Lukas Cyganek

Göttingen, November 2012