Plasticity-related gene 5 induces spine formation in immature primary neurons.

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"Back in the 1960s and early '70s I took plenty of LSD. A lot of people were doing that in Berkeley back then. And I found it to be a mind-opening experience. It was certainly much more important than any courses I ever took."

Kerry Mullis
I would like to express my deepest and sincere gratitude to Frau Juniorprof. Anja Bräuer for her supervision, guidance and the trust shown me in recent years.

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I’d like to dedicate my thesis to Riccardo, Anette, Alessandro e Rakel
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**Abbreviations**

aa  Amino acids
AMPA  α-amino-3-hydroxyl-5-methyl-4-isoxazole-propionate
ATX  Autotaxin
bHLH  Basic helix-loop-helix
CA-1  Cornus ammonis-1
C1 – C3  Consensus phosphatase sequence
C1P  Ceramid-1-phosphate
CNS  Central nervous system
DIV  Days in vitro
dNTPs  Deoxynucleotide Triphosphate
DPE  Downstream promoter element
DTT  Dithiothreitol
E  Embryonal day
EGFP  Green fluorescence protein
GAPDH  Glyceraldehyde-3-phosphate dehydrogenase
GFAP  Glial fibrillary acid protein
Glu  Glutamate
GTFs  General Transcription Factors
h  Human
HPRT  Hypoxanthine-guanine phosphoribosyltransferase
Inr  Initiator sequence
IB  Immunoblot
Ipa1  Importin alpha 1
LPA  Lysophosphatidic acid
LPC  Lysophosphatidylcholine
LPP  Lipid phosphate phosphatase
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>LPRs</td>
<td>Lipidphosphatase-related proteins</td>
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<td>LTP</td>
<td>Long-term Potentiation</td>
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<tr>
<td>m</td>
<td>mouse</td>
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<td>MAG</td>
<td>Monoacylglycerol</td>
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<td>NMDA</td>
<td>N-methyl-D-aspartic acid</td>
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<tr>
<td>P</td>
<td>post-natal day</td>
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<tr>
<td>PA</td>
<td>Phospholipase A₂</td>
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<tr>
<td>PAP2D</td>
<td>Phosphatidic acid phosphatase type 2D</td>
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<tr>
<td>PBS</td>
<td>Phosphate buffer</td>
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<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
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<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
</tr>
<tr>
<td>PIC</td>
<td>Preinitiation complex</td>
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<tr>
<td>PLD</td>
<td>Lysophospholipase D</td>
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<td>PRGs</td>
<td>Plasticity-related genes</td>
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<tr>
<td>r</td>
<td>Rat</td>
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<tr>
<td>RLM</td>
<td>RNA ligase-mediated</td>
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<tr>
<td>S1P</td>
<td>Sphingosin-1-phosphate</td>
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<tr>
<td>TBP</td>
<td>Transcription Binding Protein</td>
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<tr>
<td>TFII</td>
<td>Transcription Factor II</td>
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<td>TSS</td>
<td>Transcription start site</td>
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<tr>
<td>YPF</td>
<td>Yellow Fluorescent Protein</td>
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<td>WT</td>
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Zusammenfassung

Die PRG-Familie, eine Gruppe von fünf Membranproteinen, wurde als wirbeltierspezifisch definiert und ist hauptsächlich im Gehirngewebe exprimiert.

Ihre biologische Funktion beinhaltet die Erweiterung der Neuriten, die axonale Wegfindung und die Reorganisation nach einer Läsion. Insbesondere die Deletion von PRG-1 führt zu einer starken Übererregbarkeit des Hippocampus, während die Überexpression von PRG-3 in HeLa- und COS7-Zellen die Anzahl der physiologischen Filopodien erhöht. Im zentralen Nervensystem ist die PRG-1-Expression auf die postsynaptischen Dendriten von glutamatergen Neuronen beschränkt.


Danach habe ich das letzte Mitglied der PRG-Familie, plasticity-related gene-5 (PRG-5) charakterisiert, das in der Lage ist, verschiedene spine-Strukturen in jungen primären Neuronen zu induzieren.


PRG-5, das neue Multi-Spanning Membranprotein, lokalisiert und fördert die spine-Induktion in jungen primären Neuronen. Die Aminosäuren innerhalb der C1-C3 Domains sind für die spine-Induktion von Mäusen bei DIV2 und DIV4 verantwortlich.
In der Tat zeigen Mutagenese-Experimente in PRG-5 Rückstände von Aminosäuren, die wichtig für die spine-Induktion sind. Unsere Daten zeigen, dass PRG-5 an der spine-Induktion in primären Neuronen beteiligt ist und damit den Umbau der Plasmamembran moduliert.
Summary

The PRG family, a set of five trans-membrane proteins, were shown to be vertebrate specific and mainly expressed in brain tissue. Their actions include neurite extension, axonal path finding and reorganization after lesion. In particular, deletion of PRG-1 results in severe hippocampal overexcitability, while overexpression of PRG-3 in HeLa and COS7 cells increases the physiological number of filopodia. In the central nervous system, PRG-1 expression is restricted to postsynaptic dendrites on glutamatergic neurons.

In this study, first of all I shortly focus my attention in the upstream region of mouse and human PRG-1; PRG-1 expression is under the control of a TATA-less promoter with multiple transcription start sites. The promoter architecture of PRG-1 resulted in the identification of 450 bp, mediating approximately 40 fold enhancement of transcription in cultured primary rat neurons, compared to controls.

Afterwards, I characterized the last member of PRG family, plasticity-related gene-5 (PRG-5) which is able to induce spine-like structure in young primary neurons.

The shape of a neuron’s dendritic arbour determines the set of axons with which it may form synaptic contacts, thus establishing connectivity within neural circuits. Dynamic cytoskeleton remodelling is an essential step during this process. Putative extracellular cues may act through membrane proteins that relay signals to a network of intracellular signalling pathways, which ultimately converge on the cytoskeleton. However, the molecular mechanisms involved in these steps are not well understood.

The novel member of the vertebrate- and brain-specific PRG family, plasticity-related gene 5 (PRG-5), a multi-spanning membrane protein, localizes to and promotes the induction of spines in young primary neurons. A set of amino acid within the C1 – C3 domains are responsible for spine formation in primary neurons at DIV 2 and DIV 4.

In fact, mutagenesis experiments in PRG-5 show residual amino acid that are important for the induction of spines. Our data show that PRG-5 may be involved in spine induction in primary neurons and thereby modulate plasma membrane rearrangement.
1. Introduction

The plasticity-related gene family (PRG), comprising integral multi-spanning membrane proteins, is a subclass of the lipid phosphate phosphatases (LPP) superfamily, which control the level of lipids in the extracellular space (27, 29, 30, 50) and the regulation of effects induced by lysophosphatidic acid (LPA) (6).

LPA is a simple lysophospholipid, a metabolite in the biosynthesis of membrane phospholipids and a well-characterized signaling molecule (37). LPA is present in biological fluids and activates cells through families of G-protein-coupled receptors. This leads to multiple biological functions including cell proliferation, membrane depolarization, and cytoskeletal remodeling in numerous cell types (37). Extracellular LPA is produced in inflammatory conditions by secretory phospholipase A\textsubscript{2} acting on PA (21). LPA is also partially derived through an extracellular lysophospholipase D (lysoPLD) from a major circulating lipid, lysophosphatidylcholine (Fig. 1.1). This lysoPLD has been identified as autotaxin, an ecto-phosphodiesterase (68).

![Fig 1.1 Metabolism of phosphatidate and lysophosphatidate. Modified from (47)](image-url)
Increasing the degradation of extracellular LPA, or attenuating signaling through the LPA receptors can regulate the extracellular actions of LPA providing a control of diverse signaling cascades.

![LPA metabolism diagram](image)

**Fig. 1.2** LPA metabolism. A major route for the synthesis of extracellular LPA is through the actions of ATX by conversion of lysophosphatidylcholine (LPC) to LPA. The role of LPA in controlling signal transduction is regulated by the ecto-enzymes lipid phosphate phosphatases (LPPs), which are responsible for its degradation to monoacylglycerol (MAG) and also for controlling signaling events downstream of receptor activation (6).

Lipid phosphate phosphatase-1 (LPP1) regulates the degradation of extracellular LPA as well as the intracellular accumulation of lipid phosphates (46). More in general, the effects of the members of the lipid phosphate phosphatases superfamily, LPP1-3 are in the direct opposition to those of ATX. LPPs show a ubiquitous expression pattern (31, 63) and the mRNA are also expressed in the brain, whereas levels of LPP-2 transcripts are somewhat lower and more restricted (27, 31, 63).
All the members of the LPP superfamily are characterized by six membrane-spanning domains with three extracellular loops (27, 29, 30, 50) and are able to dephosphorylate LPA to mono-acyl-glycerol (MAG) (Fig. 1.2) (9, 10, 74). The conserved enzymatic active domains are localized within the three extracellular loops. The conserved amino acids, essential for an ecto phosphatase activity, are shown in Tab 1 (74). LPP-1 is partially expressed on the plasma membrane of cells with the C- and N-terminal inside the cell (30) and a glycosylation site on an hydrophilic loop between the first and second active site domains (9, 49, 57). LPP-1, LLP-1a and LPP-3 are partly located in the plasma membrane of some cells (29, 30, 39, 53). Their observed ability of the LPPs to dephosphorylate exogenous lipid phosphates (29, 30) implies that the LPPs could regulate circulating concentrations of LPA or S1P in biological fluids.

1.1 Plasticity-related genes

Recently, a family of plasticity-related genes (PRGs), or LPP-related proteins (LPRs) that show structural similarity to the LPPs has been identified (Fig. 1.3). These proteins, like LPPs, all contain six putative transmembrane regions. Unlike other members of the LPP family, PRG-1 and PRG-2 (LPR-4 and -3, respectively) have very long hydrophilic C-terminal tails.

**Fig. 1.3** Phylogenetic tree between mouse LPPs and mouse plasticity-related genes (PRGs) calculated using the Clustal W algorithm (6).
Mouse PRG-1 (GeneBank Accession no. AAP41099) protein contains 766 aa, PRG-2 (NP_859009) 746 aa, while PRG-3 (NP_848871), PRG-4 (Q8VCY8) and PRG-5 (AAS80161) are 325 aa, 343 aa and 321 aa, respectively. Although all PRGs protein show a high identity to LPP-family members, included the same six spanning membrane structure, they lack the residues responsible for a phospho enzymatic activity (Tab 1) (26, 61, 64, 74). Nevertheless, PRGs are involved in a variety of neuronal processes like filopodia formation, neurite extension, axonal path finding and reorganization after lesions (7, 43, 51, 58).

<table>
<thead>
<tr>
<th></th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
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<tbody>
<tr>
<td>mPRG-1</td>
<td>EFILTVCKP</td>
<td>SQH</td>
<td>TRITCYKHPVSD</td>
</tr>
<tr>
<td>mPRG-2</td>
<td>EFILTVCKP</td>
<td>SQH</td>
<td>TQITQYRHPVSD</td>
</tr>
<tr>
<td>mPRG-3</td>
<td>EFILTVCKP</td>
<td>SQH</td>
<td>NRVAEYRNMWD</td>
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<td>mPRG-4</td>
<td>EFILTVCKP</td>
<td>CRD</td>
<td>VEVAEYRNMWD</td>
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<td>mPRG-5</td>
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<td>SKE</td>
<td>NRVAEYRNMWD</td>
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<tr>
<td>mLPP-1</td>
<td>EFIALCNP</td>
<td>SQH</td>
<td>SRYSDYKHVS</td>
</tr>
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</table>

Tab. 1 The consensus phosphatase sequence motif is shown in alignment between PRGs and LPP-1 of mouse (58).

In details, through my thesis, I investigated two distinct aspect of a gene regulation: the upstream genomic sequence region of PRG-1, indicated as a putative “proximal promoter” and may be responsible of neuron specific expression of PRG-1.

Then, I characterized the new member of PRG family, PRG-5. The overexpression of PRG-5 is able to induce spine structure in young primary neurons.

1.2 Plasticity-related gene 1 (PRG-1)

PRG-1, the first member of the PRG-family, was discovered in a screen to identify proteins involved in reorganization in the dentate gyrus, following entorhinal cortex lesion, which is an
established model for the analysis of lesion-induced plasticity (52, 59). The six spanning membrane structure is showed in Fig. 1.4.

The mRNA of PRG-1, prominently expressed in the CNS and vertebrate-specific, was detected as early at embryonic day 19 (E19) in the subventricular zone and hippocampal anlage. After birth, PRG-1 transcripts are strongly upregulated in the hippocampus and entorhinal cortex, and the protein is thus found exclusively on glutamatergic neurons in the CNS (6).

![Structure model based on the amino acid sequence of mouse PRG-1. PRG-1 contains six putative transmembrane regions and the C- and the N-terminus located intracellularly. Modified from (7).](image)

**Fig. 1.4** Structure model based on the amino acid sequence of mouse PRG-1. PRG-1 contains six putative transmembrane regions and the C- and the N-terminus located intracellularly. Modified from (7).

Analysis of PRG-1 protein during excitatory synaptic transmission is thought to depend on an interaction of PRG-1 and lipid phosphates like LPA, acting through presynaptic LPA2-receptors (66).
1.2.1 Upstream region of PRG-1: a “proximal promoter”

Transcription of a eukaryotic protein-coding gene is preceded by multiple events; these include decondensation of the locus, nucleosome remodeling, histone modifications, binding of transcriptional activators and coactivators to enhancers and promoters, and recruitment of the basal transcription machinery to the core promoter. Core promoter elements define the site for assembly of the transcription preinitiation complex (PIC) and include TATA sequence, located upstream of the transcription start site (TSS), and an initiator sequence (Inr), encompassing the start site. Promoters can include a TATA box, an Inr sequence, or both of these control elements. A third core element, the downstream promoter element (DPE), was initially described in *Drosophila* and is located about 30 bp downstream of the start site (12). The DPE appears to function, in conjunction with the Inr element, as a TFIID binding site at TATA-less promoters. The core promoter includes also DNA elements that can extend ~35 bp upstream and/or downstream of the transcription initiation site (TSS). Most core promoter elements appear to interact directly with components of the basal transcription machinery. The basal machinery can be defined as the factors, including RNA polymerase II itself, that are minimally essential for transcription in vitro from an isolated core promoter (24, 41, 70) (Fig. 1.5).

**Fig. 1.5** Schematic depiction of the transcription preinitiation complex (PIC). PIC assembly is nucleated by TBP binding to the TATA box, inducing a sharp bend in the DNA template, followed by association of TFIIB, RNA pol II/TFIIF, TFIIE, and TFIIH. Each pattern denotes a distinct general transcription factor. Subunit composition is indicated, except for TFIIH (9 subunits) and RNA pol II (12 subunits). Although PIC assembly can occur by stepwise addition of the general transcription factors (GTFs) in vitro, the discovery of RNA pol II holoenzyme complexes that include GTFs suggests that stepwise assembly might not occur in vivo (24).
Although core promoters for RNA polymerase II were originally thought to be invariant, they have been found to possess considerable structural and functional diversity (60). Furthermore, it appears that core promoter diversity makes an important contribution to the combinatorial regulation of gene expression (60).

PRG-1 was found exclusively on glutamatergic neurons in the CNS. Analysis of PRG-1 null mouse models elucidated the modulatory role of the postsynaptically localized PRG-1 protein during excitatory synaptic transmission on glutamatergic neurons. This novel molecular mechanism of adaptation in neuronal transmission is thought to depend on an interaction of PRG-1 and lipid phosphates like LPA, acting through presynaptic LPA2-receptors (66).

1.3 Plasticity-related gene 5

PRG-5, previously described as phosphatidic acid phosphatase type 2D (PAP2D) (62), was recently found, like PRG-3 (58), to increase the number of filopodia in N1E-115 neuronal cells (11). PRG-5 is located inside the same chromosome of PRG-1 (chromosome 4 in human and chromosome 3 in mouse), consecutively and in an opposite orientation (Fig. 1.6) and it shows the same six spanning membrane structure, with intracellular N- and C- terminus. Overexpression studies will be performed to show the capacity of PRG-5 to induce spine structure in immature primary neurons.

![Fig. 1.6 Schematic model of chromosome 3 in mouse, with opposite orientation of PRG-1 and PRG-5.](image)
1.3.1 Spines formation and the hypothetical role of mouse PRG-5

Neuronal activity comprises an enormous number of contacts between somata, axons and dendrites, known collectively as synapses. Synapses regulate the electric communication within neural networks and most excitatory synapses in the mammalian brain are formed from tiny dendritic protrusions, named dendritic spines (4) (Fig. 1.7).

![Diagram of presynaptic bouton and postsynaptic spine](image)

**Fig. 1.7** (a) A CA-1 pyramidal cell from a hippocampal slice. The cell is filled with Lucifer Yellow, scanned and reconstructed by a confocal laser scanning microscope. (b) Example of dendritic segment with spines, selected from the cell in (a). (c) Diagram of the presynaptic bouton and a postsynaptic spine. The spine head carries three classes of glutamate receptors: the metabotropic glutamate receptor (mGluR), the ionotropic AMPA receptor (AMPA-R) and the ionotropic NMDA receptor (NMDA-R). The bouton contains transmitter vesicles with glutamate (glu). The red squares represent proteins carried by a lipid vesicular transport system to support membrane expansion and functional adjustment of spines (1).

Five main morphological stages can be identified in dissociated cultures of hippocampal neurons (16) (Fig. 1.8). Shortly after the cells attach to the substrate, motile lamellipodia develop around...
the periphery of the cell (stage 1). The significance of lamellipodia in neuronal development is unclear (16). At stage 2, development is characterized by the transformation of lamellipodia into distinct processes, which, in the course of a few hours, extend to a length of 10-15 µm (56). Several hours after the appearance of minor processes, a rather abrupt change occurs: one of the minor processes begins to grow at a much more rapid rate. From this stage onward (stage 3), its rate of growth will average 5-10 times greater than the other processes of the cell. This process is the axon. At stage 4, like the axon, dendrites develop from the minor processes that appear during the first day in culture, but significant dendritic growth begins only after about 4 d in culture, 2-3 days after axonal outgrowth (16).

Fig. 1.8 Schematic representation of neuronal polarization in cultured rat embryonic hippocampal neurons. Shortly after plating, the neurons form small protrusion veils and a few spikes (stage 1). These truncated protrusions have growth cones at their tips, and develop into several immature neurites (stage 2). One neurite then starts to break the initial morphological symmetry, growing at a rapid rate, and immediately establishing the polarity (stage 3). A few days later, the remaining neurites elongate and acquire the characteristics of dendrites (stage 4). Approximately seven days after plating, neurons form synaptic contacts through dendritic spines and axon terminals, and establish a neuronal network (stage 5) (16).

Despite different knowledge of dendritic spine were described, key aspects remain unclear, as “how do spines form?” Several lines of evidence suggest that dendrites produce long and thin dendritic filopodia that exhibit dynamic growth, allowing them to capture some of the nearby axons. (16, 54) (Fig. 1.9). Choosing and capturing the appropriate presynaptic axon via activity-
dependent or -independent signaling stabilizes the contact and the filopodia mature into dendritic spines. Filopodia grow and retract within minutes or even seconds and their motility is regulated by synaptic activity (3, 17, 33, 48, 69, 73).

**Fig. 1.9** Morphology of dendritic protrusions, filopodia and spines. Dendrites of GFP-transfected hippocampal neurons cultured for 7 days (d.i.v. 7) and 21 days (d.i.v. 21). At the immature stage (7 days), dendritic protrusions are very thin and long; these protrusions are called dendritic filopodia. In contrast, at the mature stage (21 days), dendrites are covered by dendritic spines, which commonly have an expanded head and a narrow neck. Scale bar, 5 mm. Schematic representation of morphologies of filopodium and three types of dendritic spine: thin type, stubby type and mushroom type. Gray disks represent the PSD structure and chains of red circles represent F-actin (54).

Furthermore, time-lapse imaging studies have found that spines can form directly from dendritic shafts rather than transitioning from filopodia (15).

Small spines can change their form rapidly (22, 42, 65), either disappearing or growing into large spines. By contrast, large spines are relatively stable *in vitro* (42) and survive for more than a month (65) or even for a year (22) in the mouse neocortex *in vivo*. However, these studies do not directly address the correlation between spine structure and function.
The in vivo imaging of YFP-expressing mice showing that dendritic filopodia are indeed highly dynamic and can transform into spines provides more evidence consistent with the role of filopodia in spinogenesis (22, 76). Normally, a mice at one month of age show ~12% of filopodia; in almost 4h, the 15% of filopodia form a bulbous head and more the 40% of these spine-like protrusions, persist more than 24h (76). These evidences provide that filopodia are spine-precursors.

Although, that result give credence to that theory, other models of spinogenesis may also occur and should be considered. Shaft synapses can precede the formation of spine synapses, during early developmental stages (19, 73). Furthermore, time-lapse imaging studies have found that spines can form directly from dendritic shafts rather than transitioning from filopodia (15). Thus, it could be possible that presynaptic axons recognize the shafts of dendrites and induce the postsynaptic cell to form a dendritic spine directly in apposition to the axonal terminal (19, 73). Moreover, spines may emerge independently of synaptic contact, acting as beacons for axonal terminals, which would then locate and recognize these preformed spines on dendrites and make targeted contacts. To support these model, it is interesting consider that in some parts of the brain, spinogenesis can occur in the absence of synaptogenesis (8, 13). Even if the prominent idea remain that spines came from a development of dynamic filopodia, it is still unclear if filopodia are always precursor of all spines or if most spines form themselves, directly from the dendritic shaft.

Recently, various studies on the components regulating spine morphology, such as long-term potentiation (LTP), induction of LTP, or motility of spines, have been found to be associated with the remodelling of actin cytoskeleton (20, 23, 25, 36). According to these studies, LTP increases the number of spines in the hippocampus (18), while overexpression of Rac1 induces changes in the morphology of spines (38). Spine-like structures from neurons expressing EGFP alone showed altered morphology after overexpression of Kalirin-7 in primary cortical neurons cultured for DIV 7, exhibiting a high number of protrusions of larger size and more complex shape than control neurons (45). Morphogenesis and maturation of dendritic spines are regulated upstream by the activation of the ephrinB-EphB receptor, which regulates the translocation of Rho-GEF Kalirin and the activation of Rac1 (44). Downregulation of the protein Septin 7, normally expressed in cultured neurons between DIV 0 and DIV 20, affects dendritic-spine morphogenesis (71) and loss of function of WAVE1 (Wiskott-Aldrich syndrome protein
(WASP)-family verprolin homologous protein 1) reduces the number of mature dendritic spines (32). WAVE1 also regulates the formation of the filamentous actin cytoskeleton by phosphorylation/dephosphorylation processes. Activation of WASP protein by cdc42 and Arp2/3 complex may induce the initiation of filopodia and thus regulate dendritic spine morphology (32).

Sigal et al. recently found that overexpression of PRG-3/LRP1 in HeLa and COS-7 cells increases the physiological number of filopodia independently of Cdc42 and ARP2/3 (58). Mainly expressed in the brain, PRG-3 transcripts were also found in liver, kidney and testis even through a much lower extend compared to PRG3 mRNA expression in the brain (51). In recent years, we have identified a further member of the PRG family, named plasticity-related gene-5 (PRG-5). Here, I showed that PRG-5 induce spine-like structure in young primary neurons.
2. The aim of the work

PRG-1 and PRG-5 are both members of the plasticity-related genes (PRGs), a new class of transmembrane protein, mainly expressed in brain tissue. In my thesis, I focused my attention in two distinct aspects of these genes.

I investigated the regulatory elements of the upstream region of PRG-1 and then, I characterized the new member of PRG family, PRG-5, able to induce spine structures in young primary neurons.

**PRG-1:** the upstream region of PRG-1, a DNA chromosomal region before the transductional start point (ATG), was amplified by PCR and cloned in a reporter expression vector, pGL-3 luciferase basic vector, to analyze the putative „proximal promoter“ responsible of the activation and neuron specific expression of PRG-1.

**PRG-5:** I analyzed the putative protein structure and the mRNA expression level of PRG-5 in different mouse tissues, cellular types and developmental stages of mouse. Afterwards, I investigated the intracellular localization of PRG-5 in neurons cultivated at least one day in vitro and its ability to induce spine-like structures in these young primary neurons.

Finally, I showed, by mutagenesis experiments, the amino acids of PRG-5 directly involved in the spine-induction process.
3. Material and Methods

3.1 Material

3.1.1 Laboratory equipment

Balance                                Mettler-Toledo GmbH, Giessen
Centrifuges                             RC-M150 GX, RC-5B, Ultra Pro80, Sorvall GmbH, Bad Hamburg
Dual Luciferase system                  FB12 Luminometer, Bethold
Filmprocessor                          Curix 60, Agfa-Gevaert, NV, Mortsel, Belgium
Gel filtration column                  Amersham Pharmacia Biotech Europe GmbH, Freiburg
Gel electrophoresis equipment           Bio-Rad Laboratories GmbH, München
GeneAmp PCR system 9600                 PE Biosystems, The Perkin Elmer Corporation, CA, USA
Heating block                          Grant Instruments, Cambridge, UK
Incubator for HEK-293 and neuron cells Memmert, Germany
Incubator shaker                       New Brunswick scientific GmbH, Nürtingen
Microscope                              Eppendorf AG, Hamburg
Confocal microscope                    Leica TCS SP5, Microsystem, USA
Mx3005P Real Time PCR System           Stratagene, La Jolla, CA, USA
Pipettes                                Eppendorf AG, Hamburg
pH meter                                Mettler-Toledo GmbH, Giessen
Spectrophotometer                       Bio-Rad Smart sectm3000, BioRad Laboratories GmbH, München
Thermocycler                            PTC100, MJ Research, Inc.; Watertown, USA Biometra, Biometra GmbH, Germany
                                       Bior-Rad Laboratories GmbH, München
Vertical gel electrophoresis system     Bio-Rad Mini Protean II
UV-Transiluminator                     Biometra, Götttingen
<table>
<thead>
<tr>
<th>Equipment</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vortex</td>
<td>Vortex Genie 2, Bender und Hobein AG, Zürich, Switzerland</td>
</tr>
<tr>
<td>Water bath</td>
<td>GFL, Germany</td>
</tr>
<tr>
<td>Western blotting apparatus</td>
<td>Bio-Rad Transblot SD apparatus</td>
</tr>
</tbody>
</table>

### 3.1.2 Solutions and buffers.

<table>
<thead>
<tr>
<th>Description</th>
<th>Composition and Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibiotics (1000x)</td>
<td>50mg/ml Ampicillin&lt;br&gt;30mg/ml Kanamycin</td>
</tr>
<tr>
<td>LB medium</td>
<td>10g Bacto-tryptone&lt;br&gt;10g NaCl&lt;br&gt;pH adjusted to 7.2; autoclaved</td>
</tr>
<tr>
<td>LB agar</td>
<td>LB medium&lt;br&gt;15g/l bacto agar</td>
</tr>
<tr>
<td>6x DNA loading buffer</td>
<td>0.2% bromophenol blue&lt;br&gt;60% Glycerol&lt;br&gt;60mM EDTA</td>
</tr>
<tr>
<td>5x RNA loading buffer</td>
<td>50% Glycerol&lt;br&gt;1mM EDTA&lt;br&gt;0.25% Bromophenol blue</td>
</tr>
<tr>
<td>50x TAE buffer</td>
<td>242g Tris base&lt;br&gt;57.1 ml Glacial acetic acid&lt;br&gt;100 ml 0.5M EDTA ph 8.0</td>
</tr>
<tr>
<td>10x TBS buffer</td>
<td>100mM tris&lt;br&gt;9% NaCl&lt;br&gt;pH adjusted to 7.4</td>
</tr>
<tr>
<td>PBS buffer</td>
<td>137mM NaCl&lt;br&gt;2.7mM KCl&lt;br&gt;4.3mM Na$_2$HPO$_4$&lt;br&gt;1.47mM KH$_2$PO$_4$&lt;br&gt;pH adjusted to 7.4</td>
</tr>
</tbody>
</table>
Blocking buffer 1x TBS
3% BSA

Western blotting running buffer 20mM Tris
150mM Glycin
20% Methanol
0.08% SDS

Ponceau-S staining solution 0.2% Ponceau
3% Trichloroacetic acid

SDS-PAGE 0.20g SDS
1.5M Tris-HCl pH 8.8
50 ml H₂O

3% Acrylamide pre-mix 3.8 ml 40% Acrylamide 29:1
0.5M Tris-HCl pH 6.9
10 mL SDS
6 ml 50% sucrose
500µl 1% Bromophenol Blue
50 ml H₂O

8% Resolving gel 2.00 ml 40% Acrylamide 29:1
1.5M Tris-HCl pH 8.8
1.70 ml SDS
1.30 ml 50% Sucrose
43.0 µl 10% Ammonium persulfate
9.0 µl TEMED
4.948 ml H₂O

Stacking gel 4.00 ml 3% Acrylamide pre-mix
28 µl 10% Ammonium persulfate
6µl TEMED (stir quickly)

2x SDS gel-loading buffer 100 mM Tris-HCl pH 6.8
4% (w/v) SDS
0.2% (w/v) bromophenol blue
20% glycerol

200 mM β-Mercaptoethanol (added just before the buffer was used)
Protein elution buffer

- 25mM Tris
- 192mM Glycine
- 0.1% SDS

Glycerol stocks plasmids clones were prepared in 25% glycerol and stored at – 80°C.

### 3.1.3 Biological substance and Kits

**Bacteria**
- DH5α chemocompetent cells, Promega, Mannheim
- XL-Gold chemocompetent cells, Stratagene, CA, USA
- BL-21 (RIL) bacterial expression cells, Stratagene, CA, USA

**HEK-293 cells**
- (ATCC, Manassas, VA, USA)

**Antibodies**
- Moleculare Probes, Invitrogen Santa Cruz
- Biotechnology Inc., CA, USA
- AbCam, USA
- Molecular Probes, Invitrogen Sigma, USA
- Sigma Aldrich - Germany

**Mice**
- C57/B16, Germany

**Enzymes and reagents**
- DNasel, RNase-free, RNaseH, Roche Diagnostics, Mannheim and Promega Madison, USA
- Taq DNA Polymerase, Promega, Madison, USA
- Taq DNA Herculase, Stratagene, La Jolla, CA, USA
- TaqMan Universal PCR Master Mix, Applied Biosystems, USA
- T4 DNA ligase, BioLabs, England
- Restriction enzymes, BioLads, England
- Ethidium bromide, Sigma, Deisenhofen
- Agarose, Roth, Karlsruhe
- Hepes, Roth, Karlsruhe
- pcDNA3.1/zeo(+) vector, Promega, Madison, USA
- pGL3basic vector, Promega, Madison, WI, USA
- pEGFP vector, Promega, Madison, USA
- Fugene6-reagent, Roche, Mannheim, Germany
Effectene, Qiagen, Hilden, Germany

Gel elution and PCR Nucleospin Extract kit, Macherey-Nagel Düren clean-up kits

Gel purification Kit Qiagen, Hilden

RT-PCR Qiagen, Hilden

TopoTA-cloning Kit Promega, Mannheim, Germany

Multi site mutation Kit QuikChange, Stratagene, CA, USA

Protein fraction PromoKine Cell Fractionation, Heidelberg, Germany

3.1.4 Software

MetaMorph Systemss 6.2r4 Universal Imaging, Downingtown, PA

Leica Confocal Software Leica

v. 2.61

Adobe Photoshop 6.0 Adobe Systems Incorporated, San Jose, CA, USA

DNASIS MAX Hitachi Software Engineering, CA, USA

ExPASy Compute pI/MW tool http://www.ic.sunysb.edu/stu/shilin/rnai.html

3.2 Methods

3.2.1 Vector construction for PRG-1

To generate promoter deletion constructs of PRG-1, I amplified specific-length fragments from the human and mouse total cDNA with Herculase (Stratagene, La Jolla, CA, USA). Amplified fragments were restricted and inserted into pGL3basic (Promega, Madison, WI, USA). Denomination of vectors referred to the position of the inserted region in relation to the translation start point, ATG.

3.2.2 Restriction enzymes

New England Biolabs (NEB) provided the restriction enzymes used in this study. The restriction enzyme digests were performed on at least 1μg DNA in 1X the final recommended buffer suggested by the manufacturer and using at least 1 unit of the specified enzymes. The reactions were incubated at 37°C for 1 hour for complete restriction enzyme digestion.

3.2.3 Agarose gel electrophoresis

Agarose concentrations ranging from 0.7% to 1% in TAE buffer (40 mM Tris-acetate, 1 mM EDTA containing 1 μg/ml ethidium bromide (EtBR; Sigma)), was used to cast agarose gels. DNA samples were loaded onto the gels with gel loading buffer (0.042% (w/v) bromophenol blue, 6.67% (w/v) sucrose) along with the DNA markers, λDNA/Pst I fragment markers (Invitrogen). Gel electrophoresis was performed between 90 to 110 V in TAE buffer for varying times to obtain optimal resolution. A gel documentation system (Bio-Rad) was used to visualize the gels by UV illumination.
3.2.4 Purifying linear DNA fragments

Linear DNA fragments were isolated from DNA agarose gels and purified using Qiagen spin columns. Based upon the fragment size or the amount of sample, electroelution was sometimes performed to elute the DNA from the gel. Elutip-D columns (Schleicher & Schuell) were used to further purify and concentrate the electroeluted DNA sample according to manufacturer’s protocol, using 200 mM NaCl, 20 mM Tris-Hydrochloric acid (HCl) and 1.0 mM EDTA, pH 7.4.

3.2.5 Isolation of plasmids: mini prep

A small scale preparation using an alkali lysis method was performed to screen colonies after transformation for positive recombinants. Single isolated colonies were selected and cultured overnight in LB containing the appropriate antibiotic. The next day, the cultured cells were resuspended in 50 mM glucose, 10 mM EDTA, 25 mM Tris- HCl (pH 8.0), 2 mg/ml lysozyme (Sigma) then lysed with a 200 mM NaOH, 1% sodium dodecyl sulfate (SDS) solution followed by the addition of 3 M sodium acetate, pH 5.2.

This precipitated out the bacteria chromosomal DNA, cellular protein and debris. The supernatant was extracted twice with phenol:chloroform:isoamyl alcohol followed by ethanol precipitation. The precipitated DNA was resuspended in TE (pH 8.0) containing 40 μg/ml RNAse A. DNA concentration was measured by standard A260/A280 spectrophotometric readings and visualized on agarose gels.

3.2.6 Bacterial cells

The bacterial host for vector propagation routinely used was DH5α E. coli cells from Promega. E. coli XL-Gold chemocompetent cells and BL-21 (RIL) bacterial expression cells were both obtained from Stratagene. These cells were grown at 37°C in a shaking incubator in a flask containing Lauria-Bertani (LB) broth consisting of 1.0% (w/v) tryptone (Difco), 0.5% (w/v) yeast extract (Difco) and 1.0% (w/v) NaCl supplemented with the appropriate antibiotic such as ampicillin (100μg/ml) or kanamycin (50 μg/ml), depending on the vector. For isolation of
transformed bacterial cells, the cells were plated onto selective LB-agar plates, containing LB broth with 1.5% (w/v) agar supplemented with the appropriate antibiotic, and incubated inverted at 37°C overnight.

3.2.7 Transient transfection & dual luciferase assay

All cells were transiently transfected with equal amounts of the various luciferase-fusion constructs or pGL3basic in serum-free growth media using transfection reagents according to the manufacturer’s protocol. pHRLtK (Promega, Madison, WI, USA) containing the Renilla luciferase gene fused to the herpes simplex virus thymidin kinase promoter was cotransfected in order to normalize transfection efficiencies. HEK-293 cells (ATCC, Manassas, VA, USA) were transfected with Fugene6-reagent (Roche, Mannheim, Germany). Primary neurons and astrocytes were transfected with Effectene (Qiagen, Hilden, Germany). The cells were lysed 48 hours after transfection and the luciferase activities of the cell lysates were determined using the dual-luciferase reporter assay system (Promega, Madison, WI, USA). Transfections with each construct were performed at least three times.

3.2.8 Statistical analysis

Statistical analysis was performed using the Mann Whitney U-test, which was performed using Prisms 5 software (Graphpad Software, San Diego, USA). Data are expressed as means +/- standard deviation. A critical value for significance of $p<0.05$ was used.

3.2.9 RNA Isolation

Organs were dissected from mice and homogenized in TRIzol reagent. Primary neurons, microglial cells, or astrocytes were scraped in 1 x PBS, centrifuged for 5 min at 900 rpm and 4°C. Cell pellets were dissolved in 1 ml TRIzol reagent. Total RNA was isolated according to the TRIzol protocol (Invitrogen). Following precipitation and drying, RNA was resuspended in RNase und DNase-free water, an aliquot quantified by $A_{260nm}$ spectrophotometry (Biomate 3 spectrophotometer, Fisher Scientific), and stored at −80°C.
3.2.10 DNAse treatment of total RNA sample and RNA cleanup

DNAse treatment was used to remove DNA from total RNA preparation. To the total RNA sample (~ 2 µg/µl diluted in RNase-free water) 5 µl 10x RQ1 DNAse buffer, 5 µl 100 µM DTT and 2 units RQ1 RNase-free DNAse (Promega) was added to a final volume of 50 µl. The mix was incubated for 30 min at 37°C. After the enzymatic reaction the volume of the sample was adjusted to 100 µl and the RNAeasy kit (QIAGEN) was used to clean up the total RNA. The RNA was eluted twice in the same tube with RNase-free water to obtain a higher total RNA amount.

3.2.11 cDNA Synthesis

cDNA was synthesized using the High-Capacity cDNA Archive Kit (Applied Biosystems) according to the manufacturer's protocol. Briefly, 5 µg total RNA was reverse transcribed in a 50 µl reaction containing 5 µl 10 x random hexamer primers, 2 µl 25 x dNTP mix and 125 U MultiScribe reverse transcriptase. cDNA was diluted 1:5 with RNase and DNase-free water and stored at −20°C. The quality of the amplified cDNA was controlled by β-actin PCR.

3.2.12 PCR

DNA was amplified by PCR using the high fidelity Pfu polymerase or Herculase enzyme. The reaction volume was 25µl.

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Template DNA</td>
<td>30 ng</td>
</tr>
<tr>
<td>10x buffer</td>
<td>5 µl</td>
</tr>
<tr>
<td>dNTP’s (10mM)</td>
<td>1 µl</td>
</tr>
<tr>
<td>Forward Primer (5 pmol)</td>
<td>1 µl</td>
</tr>
<tr>
<td>Reverse Primer (5 pmol)</td>
<td>1 µl</td>
</tr>
<tr>
<td>Pfu polymerase (3 U/µl)</td>
<td>0.5 µl</td>
</tr>
<tr>
<td>H2O</td>
<td>25 µl</td>
</tr>
</tbody>
</table>

The following PCR profile was used:

PRG-5-Flag
1. Initial denaturation  95°C  2 min
2. Denaturation  95°C  45 s
3. Annealing  64°C  30 s
4. Extension  72°C  1 min for every 1000 bps
5. 30 cycles
6. Final extension  72°C  10 min

pGL-3-PRG-1

1. Initial denaturation  95°C  5 min
2. Denaturation  95°C  1 min
3. Annealing  66°C  45 s
4. Extension  72°C  1 min for every 1000 bps
5. 30 cycles
6. Final extension  72°C  10 min

The PCR products were agarose gel extracted.

3.2.13 Molecular cloning of PRG-5 and sequence analysis

Vector construction of pcDNA3.1Zeo(+)/PRG-Flag involved ligation of the gel-purified 0.9 kb of PRG-5-Flag fragment (EcoI/NotI cut site) into the pcDNA3.1Zeo(+) vector, resulting in the 6.4 kb vector (Fig. 2.1).

PRG-5 full-length clones were amplified by reverse transcription-polymerase chain reaction from postnatal mouse cDNA. Database searches used BLAST on the website of the National Center for Biotechnology Information. Multiple sequence alignments (GenBank Accession no. ACJ60628, human PRG-5; AAS80161, mouse PRG-5; NP_001101190, rat PRG-5) were made using DNAsis Max v2.0 software (Hitachi, Olivet Cedex, France). Transmembrane structure was predicted with ProDom and Swissprot databases. cDNA construct of mouse PRG-5, containing a C-terminal Flag, were used for the transiently transfection of HEK-293 cells and primary neurons cells. Briefly, cDNA from mouse brain was amplified by PCR using primers:

For_EcoRI: 5’ – gaa ttc atg ccc ctg ctg ccc gtg gcg ctc atc agc – 3’
Rev_NotI: 5’ – gcg gcc gct cag tcg tca tcg tct ttg tag tct gtg act
tcc gca aag gca gtg aeg tgg ttc tgc ag – 3’
Reverse primer contains the Flag sequence. The oligonucleotides were synthesized by Metabion (Munich, Germany). The PCR products were purified by column (Qiagen), sub-clone in TOPA-Ta vector (Invistrogen), checked the sequence and then cloned in pCDNA3.1(+)zeo expression vector (Invitrogen) by EcoRI and NotI.

3.2.14 Ligation and transformation

Purified PCR products were cloned in a total volume of 25 µl ligation reaction as below. This reaction mix was incubated at room temperature for 1 hour.

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X buffer</td>
<td>2.5</td>
<td>µl</td>
</tr>
<tr>
<td>Vector</td>
<td>5</td>
<td>µl</td>
</tr>
<tr>
<td>Insert</td>
<td>15</td>
<td>µl</td>
</tr>
<tr>
<td>T4 - Ligase</td>
<td>1</td>
<td>µl</td>
</tr>
<tr>
<td>H₂O</td>
<td>1.5</td>
<td>µl</td>
</tr>
</tbody>
</table>

Fig. 2.1 Diagram of pcDNA3.1Zeo(+) vector
For transformation, 16 µl of the ligation reaction were added to 750 µl thawed competent bacteria and incubated on ice for 10 minutes. Then, bacteria were heat shocked for 30 seconds at 42°C and put on ice 2 minutes. After that, 750 µl SOC media was added and incubated at 37°C for 1 hour with 800 rpm shaking.

### 3.2.15 Clone analysis

Extraction of the plasmids of transformed bacterial cells was carried out by Minipreps. The restriction enzyme digestion reaction mix contained:

- 10X buffer: 2µl
- Restriction enzyme (10U/µl): 1µl
- DNA: 1µl
- H₂O: 16µl

Bacterial clones shown to contain the target plasmids were preserved for further use. Sterile glycerol was added in a ratio of 1:3 and aliquoted in 1.5 ml cryotubes then stored at –80°C.

### 3.2.16 Site direct mutants of mouse PRG-5

We have synthesised two complimentary oligonucleotides, containing the nucleotides we had mutated (uppercase) and generated mutants of mouse PRG-5 using the QuikChange protocol (Stratagene)

<table>
<thead>
<tr>
<th>Oligonucleotide</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>F_Mut_S193W</td>
<td>5’ – cgg aaa aca ttc cca TGT aag gaa gct gcc ctg – 3’</td>
</tr>
<tr>
<td>R_Mut_S193W</td>
<td>5’ – cag ggc agc ttc ctt CCA tgg gaa tgt ttt ceg – 3’</td>
</tr>
<tr>
<td>F_Mut_E195G</td>
<td>5’ – aca ttc cca tcc aag GGC gct gcc ctg agt gtc – 3’</td>
</tr>
<tr>
<td>R_Mut_E195G</td>
<td>5’ – gac act cag ggc age GCC ctt gga tgg gaa tgt – 3’</td>
</tr>
<tr>
<td>F_Mut_E195H</td>
<td>5’ – aca ttc cca tcc aag CAC gct gcc ctg agt gtc – 3’</td>
</tr>
<tr>
<td>R_Mut_E195H</td>
<td>5’ - gac act cag ggc age GTG ctt gga tgg gaa tgt – 3’</td>
</tr>
<tr>
<td>F_Mut_R241E</td>
<td>5’ - ctt act gga ctc aac GAG gta geg gaa tat cga – 3’</td>
</tr>
<tr>
<td>R_Mut_R241E</td>
<td>5’ - tcg ata ttc cgc tac CTC gtt gag tcc agt aag – 3’</td>
</tr>
</tbody>
</table>
Briefly, PRG-5 was amplified by PCR using the high fidelity *Pfu* Turbo DNA and mutated primers in a final reaction volume of 50μl:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Template DNA</td>
<td>30 ng</td>
</tr>
<tr>
<td>10x buffer</td>
<td>5 µl</td>
</tr>
<tr>
<td>dNTP’s (10mM)</td>
<td>1 µl</td>
</tr>
<tr>
<td>F_Mut Primer (125 ng)</td>
<td>1 µl</td>
</tr>
<tr>
<td>R_Mut Primer (125ng)</td>
<td>1 µl</td>
</tr>
<tr>
<td><em>Pfu</em> polymerase (2,5 U/µl)</td>
<td>1 µl</td>
</tr>
<tr>
<td>H2O</td>
<td>38 µl</td>
</tr>
</tbody>
</table>

The following PCR profile was used:

1. Initial denaturation 95°C 30 s
2. Denaturation 95°C 45 s
3. Annealing 55°C 1 min
4. Extension 68°C 1 min for every 1000 bp
5. 18 cycles
6. Storage 4°C hold

We added 1 µl of the *Dpn* I restriction enzyme (10 U/µl) directly to each amplification reaction and incubated each reaction at 37°C for 1 hour to digest the parental (i.e., the nonmutated) supercoiled dsDNA before transfection in E.Coli.

### 3.2.17 HEK-293, preparation and transfection

HEK-293 cells, containing the adenoviral E1 genes transformed into the cells, were purchased from Microbix (ATCC, Manassas, VA, USA) and grown in Minimum Essential Medium containing Earle’s Salts (EMEM; Gibco) supplemented with 10% FBS and gentamicin solution. The cell lines were mostly passaged using Trypsin/ethylenediamine tetracetic acid (EDTA; Gibco) whereas HEK-293 cells were passaged using a 1X citric saline solution (10% (w/v) KCl and 4.4% (w/v) sodium citrate). All cells lines were cultured in a 37°C humidified CO2 incubator with a 5% CO2 atmosphere. Trypan-Blue (Gibco) was used for cell counting on a haemocytometer.

HEK-293 cells were routinely maintained at 37°C with 5% CO2 in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (Gibco), 100 U/mL penicillin and
100µg/mL streptomycin, respectively as previously described (7) into 12-well plates or dishes with cell density of 10,000 – 20,000 cells/cm² and transfected with calcium; in two separate tubes I prepared; solution A:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA</td>
<td>1-2 µg</td>
</tr>
<tr>
<td>CaCl₂ 2M</td>
<td>12.3 µl</td>
</tr>
<tr>
<td>H₂O to</td>
<td>100 µl</td>
</tr>
</tbody>
</table>

solution B:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBS 2x</td>
<td>100µl</td>
</tr>
</tbody>
</table>

Add the 100µl solution A to the solution B (100 µl) and vortex for a few seconds. Incubate at RT for a 3-5 min, mix again and dropwise onto the media.

### 3.2.18 Primary neurons, preparation and transfection

Primary neurons were isolated from C57BL/6 mouse embryos at embryonic day 18. Brain from several embryos were collected and washed twice in ice-cold HBSS (Invitrogen). The tissue was incubated in 4 ml HBSS and 400µl trypsin (Invitrogen) for 15 min at 37°C, resuspended in plating medium (MEM supplemented with 10% horse serum, Invitrogen, 0.6% glucose, 100 U/ml penicillin and 100µg/ml streptomycin). After dissociation, the neurons were plated in plating medium onto poly-L-lysine (Sigma)-coated cover slips into 12-well plates with cell density of 100.000 cells/cm². Three hours after plating, cells were washed twice with phosphate-duffered saline (1 x PBS) and incubated in Neurobasal A medium (Invitrogen) supplemented with 2% B27 (Invitrogen), 0.5 mM glutamine, 100 U/ml penicillin, 100 µg/ml streptomycin at 37°C and 5% CO₂. Primary neurons were routinely maintained at 37°C with 5% CO₂ in Neurobasal A medium (Invitrogen) supplemented as described above for 1 or 3 days in vitro and transfected using efectine (Qiagen).
3.2.19 qRT-PCR

Each PCR reaction contained: 8 µl H₂O, 10 µl TaqMan® Universal PCR Master Mix (Applied Biosystems), 1 µl cDNA and 1 µl TaqMan Gene Expression Assays: for PRG5 (Assay ID Mm01310525_m1), GAPDH (glyceraldehyde-3-phosphate Dehydrogenase; Assay ID 4352932E) and β-actin (Assay ID 4352933E). For HPRT (hypoxanthine phosphoribosyltransferase), separate primer and probe were used primer mix:

For: 5’ – atc att atg ccg agg att tgg aa – 3’
Rev: 5’ – ttg agc aca cag agg gcc a – 3’

And probe:
5’ – tgg aca gga ctg aaa gac ttc gag atg – 3’

The reactions were run on the ABI PRISM™ 7700 Sequence Detection System (Applied Biosystems). Standard curves were produced with serial dilutions of cDNA from P5 mouse brain with amplification efficiency between 90 and 100%. Each result is the average of at least three separate experiments.

3.2.20 Immunocytochemistry

Primary neurons and HEK-293 cells, were fixed, after 24 hour from the transfection, in 4% paraformaldehyde for 20 min and permeabilized with 0.2% Triton X-100 (Sigma) in PBS for 5 min, washed three times with PBS 1X and incubate with block-solution, containing 10% FCS in PBS, over night at 4°C. Cells were incubated with first antibodies 2 hours at room temperature and washed 3 times with PBS 1X. Second antibodies applied 1 hour at room temperature. After 3 more washes with PBS 1X, glasses were mounted in FluoroSave Reagent. Confocal images were processed with Leica software.
3.2.21 In situ hybridization

For hybridization, I used an antisense oligonucleotides:

\[ 5' – ggc\, agt\, gtt\, ctt\, cca\, agg\, gct\, ttc\, tact\, cg\, aat\, gtt\, gg – 3' \]

Complementary to the mouse sequence of PRG-5, the oligonucleotide was synthesized by Metabion (Munich, Germany). The specificity was confirmed by a BLAST GenBank search to rule out crosshybridization with other genes. Horizontal cryostat sections (20µm) were fixed in 4% paraformaldehyde (PFA), washed in 0.1M phosphate buffered saline (PB) (pH7.4) and dehydrated through an ascending series of pure ethanol. The oligonucleotides were end-labelled using terminal desoxynucleotide transferase (Boehringer Mannheim, Germany) and x-dATP (DuPont NEN). Probe labelling was performed for 10 min at 37°C. The radioactive probes were purified using BioSpin6 Chromatography Columns (Bio-Rad). I used 100 000-200 000 cpm labelled oligonucleotides in 50µl hybridization buffer (50% formamide, 10mM Tris-HCl, pH 8.0, 10 mM PB, pH 7.2, 2x SSC, 5mM EDTA, pH 8.0, 10% dextran sulphate, 10 mM dithiothreitol (DTT), 1mM β-mercaptoethanol and 200 ng/µl tRNA) per section. Hybridization was performed for 16h at 42°C in a humidified chamber, after which the slides were washed as follows: 1 x 60 min in 0.1 x SSC at 56°C and 1 x 5 min in 0.05 x SSC at room temperature. Finally, the sections were rinsed in H₂O at room temperature and dehydrated in 50, 75 and 96% ethanol. For autoradiography, slides were exposed to Kodak X-OMAT AR X-ray films (Amersham, Heidelberg, Germany) for 15 days. No signals were detected on sections hybridized with specific antisense oligonucleotide when unlabelled oligonucleotide was added in 100-fold surplus or with sense probes of the respective oligonucleotides. Following exposure, sections were rehydrated using a decreasing ethanol series, washed in phosphate-buffered saline, Nissl counterstained (cresyl violet acetate; Sigma, Germany), dehydrated through an ascending series of ethanol, flat-embedded with Entellan (Merck-Germany) and coverslipped. Sections were then digitally photographed (Zeiss, Axioplan, Germany).
3.2.22 SDS-polyacrylamide-gel electrophoresis (Protein)

Protein samples (prepared in Laemmli buffer followed by boiling at 95°C for 5 min) were separated on 10-12% mini SDS-PAGE gels (Mini-protein II Dual Slab Cell, Bio-Rad; Laemmli 30 3. Methods 1970). In this system, proteins denatured in the presence of SDS and 2-mercaptoethanol as thiol reducing agent acquired a rod-like shape and a uniform charge-to-mass ratio proportional to their molecular weights. The gels were stained with colloidal Coomassie stain and the protein sizes were determined by comparing the migration of the protein band to a molecular mass standard (PageRuler prestained Marker, Fermentas).

3.2.23 Western blot

Protein concentration was determined by BCA protein assay (Pierce, IL, USA) using BAS as a standard. Samples were separated by 10% SDS-PAGE and the proteins were transferred to a PVDF membrane (Amersham Biosciences, CA, USA). The membrane was incubated with primary antibody PAP2D (AbCam, UK) 1:700 or Flag antibody (Sigma) 1:1000, followed by horseradish-peroxidase-conjugated secondary antibody. The immunoreactive bands were detected with SuperSignal West Pico chemiluminescent substrate (Pierce). Subcellular protein fractions (cytosol and membrane) were prepared using the kit PromoKine Cell Fractionation Kit protocol (PromoKine, Germany).
4. Results

4.1 Identification of proximal promoter of PRG-1.

In order to elucidate the peculiar expression profile of PRG-1 in neurons, a reporter mouse model expressing YPF controlled by the native PRG-1 promoter, was studied (Geist et al., in progress). PRG-1 was able to show that 200 kbp sequence enclosing the PRG-1 gene is effectual to confer cell type specific expression. By RLM-Race in rat as well as in mouse, and confirmed these via PCR-analysis, it were mapped additional transcription start points (TSS) (Geist et al., in progress).

In relation to the translation start point ATG = +1, PRG-1 mapped several transcription start sites (TSS) in rat and mouse. Rat transcripts start at position: -201, -160, -156, -96, +46, +69 while mouse transcripts start at position: -149, -144, -101, +2, +43, +45, +71 and +79 (Geist et al., in progress) (Fig. 4.1).

![Fig. 4.1](image_url)

**Fig. 4.1** PRG-1 is transcribed from multiple transcription start points in rodents. (A) Mapped transcription start sites of PRG-1 in rat (filled arrows): -201, -160, -156, -96, +46 and +69. The positions are designated in relation to the first translation start site (ATG=+1). Mapped transcription start sites in mouse (open arrows): -149, -144, -101, +2, +43, +45, +71 and +79. A second putative translation start site is conserved in mammals, in mouse and rat at position +145, in the human sequence at position +143. (B) Position of amplificates in relation to the first murine ATG. (C) Positive control amplified from mouse genomic DNA (gDNA) for comparison of amplification efficacy. PCR products were amplified from mouse cDNA.
Some TSS were located downstream of the first translation start point, revealing a second putative translation start site, which is conserved in the mammalian PRG-1 sequences. In mouse and rat, the second ATG was located at position +148, and in human at position +145.

To further investigate the core promoter of mouse PRG-1, I performed dual-luciferase-assays with several deletion-constructs. The mouse 1869-bp fragment (m-1724/+145) was subcloned into the promoter-less luciferase reporter vector, pGL3, in sense and antisense orientations, and its promoter activity was analyzed in primary rat neurons and astrocytes. As shown in Fig. 4.2, luciferase activity was minimal, in both primary neurons and astrocytes, when expression was driven by the 1869-bp sense and antisense fragment respectively (m-1724/+145 and m-1724/+145rev), compared with the pGL3 vector.

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**Fig. 4.2** Identification of a sequence element in the mouse PRG-1 promoter sufficient to confer neuronal transcription. PRG-1 promoter 5’ end deletion constructs were ligated in the dual luciferase reporter gene vector pGL3. constructs and expression control pHRLtk (promega) were co-transfected in primary rat neurons and astrocytes. The relative luciferase activity is defined as ratio of Firefly luciferase versus Renilla luciferase. Basal-level luciferase activity of cells transfected with pGL basic and pHRLtk (promega) was taken as 1%. Results are expressed as fold increase +/-SD. The experiments were repeated at least three times independently. The positions of the constructs were designated in relation to the first translation start site (ATG=+1).
Luciferase activity was 40 fold higher in neurons, respect astrocytes, when expression was driven by the 400-bp fragment (m-277/+145) (Fig. 4.2).

Transfection of human PRG-1 promoter deletion-constructs in primary rat neurons and astrocytes resulted, as well in an identification of a 450 bp fragment (h-300/+143). The luciferase activity was 50 fold greater than that of control plasmid in neurons, compared to astrocytes (Fig. 4.3). Truncation of this fragment to 300 bp (h-169/+143) leaded to a 50% reduction of specificity for neuronal transcription down to the level of stimulation in astrocytes. Transfection of this 300 bp-sequence in reverse orientation (h+143/-169) leaded to almost complete loss of transcriptional activity. All constructs with a 3' -extension to the second putative translation start site display enhanced transcriptional activity compared to controls. Interestingly, the addition of 5'-sequences to the 450 bp minimal promoter led to a strong decrease of transcriptional activity (Fig. 4.3).

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**Fig. 4.3** Human PRG-1/pGL3 reporter constructs and expression control pHRLtk were co-transfected in primary mouse neurons and astrocytes.
4.2 Expression pattern of PRG-5

Human PRG-5 was firstly cloned as phosphatidic acid phosphatase type 2D (PAP2D) and the two isoforms were derived as PAP2D_v1 and PAP2D_v2 (62). The two isoforms differ in 15 bp at the C-terminus and the five amino acids not presented in PAP2D_v2 are VTSVQ, corresponding, in mouse PRG-5, to the amino acids VTSLQ (Fig. 4.4).

Fig. 4.4 Mouse PRG-5 nucleotide and amino acid sequence. Start and stop codons of mPRG-5 (GeneBank Accession no. AAS80161) are in red. Putative transmembrane domains (TM) are boxed in grey and were predicted with ProDom and Swissprot databases. The five amino acids not presented in PAP2D_v2 are underlined.
In my studies, I specifically investigated the longest isoform in mouse. Mouse PRG-5 full length clones were amplified by reverse transcription polymerase chain reaction from postnatal and adult mouse hippocampus cDNA. The PCR product it’s been subcloned in TA vector to check the correct sequence and then fused on the pCDNA3.1+vector (Fig. 4.5).

![Image](image-url)

**Fig. 4.5** PCR from total amount of mouse cDNA. M = Lambda marker digested by PstI. The single band presence in the lines 1 and 2 are PRG-5 amplified by PCR, using deceries amount of mouse cDNA as template. Line 3 is a negative control.

![Structure Model](structure-url)

**Fig. 4.6** Structure model based on the amino acid sequence of mouse PRG-5 (ExPasy). The double line represent the plasma membrane.
The protein contains 321 amino acids and its molecular weight measures approximately 35 KDa (Fig. 4.6). In silico analysis (ProDom and Swiss-Prot databases) of human, rat, and mouse PRG-5 revealed protein homologies conserved up to 90% (Fig. 4.7).

**Fig. 4.7** Human, mouse, and rat PRG-5 sequence alignment. Full-length human and rat sequences were taken from GeneBank. The accession numbers are AAS80161 (mouse PRG-5), ACJ60628 (human PRG-5) and NP_001101190 (rat PRG-5). Identical sequences between all three different species are shaded in yellow and between two of three changes in green.
As a PRG/LRP family member, PRG-5 is a six membrane-spanning protein with a C- and N-terminus locked on the cytoplasmic site. Although sequence alignments between all PRGs/LRPs and LPP-1 revealed conserved amino acids, especially in the catalytic sequence motif within the three loops (Tab. 1), PRG-5 lacks several crucial residues for ecto-phosphatase activity (26, 61, 64, 74).

<table>
<thead>
<tr>
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<th>C1</th>
<th>C2</th>
<th>C3</th>
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<tbody>
<tr>
<td>mPRG-1</td>
<td>(206) PFLTVCKP</td>
<td>(251) SQH</td>
<td>(297) TRITQYKRHVD</td>
</tr>
<tr>
<td>mPRG-2</td>
<td>(156) PFFTVCKP</td>
<td>(203) SQH</td>
<td>(249) TQITQYRSHFD</td>
</tr>
<tr>
<td>mPRG-3</td>
<td>(154) TFLTVQDP</td>
<td>(196) SKH</td>
<td>(235) NEVAEYRHSVSD</td>
</tr>
<tr>
<td>mPRG-4</td>
<td>(156) PHFLVCKP</td>
<td>(207) CKD</td>
<td>(254) VREVAEYRHSVSD</td>
</tr>
<tr>
<td>mPRG-5</td>
<td>(149) PHFLAICP</td>
<td>(193) SKE</td>
<td>(240) NEVAEYRHSVSD</td>
</tr>
<tr>
<td>mLPP-1</td>
<td>(126) PHFLAICP</td>
<td>(169) SGH</td>
<td>(216) SVSRYKMRHVD</td>
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**Tab. 1** The consensus phosphatase sequence motif between PRGs and LPP-1 of mouse. In red the crucial amino acids of LPP-1 (58).

Through northern blot analysis with Pan Oligo, it is known that PRG-5 is mainly expressed in the brain and weakly in testis (Fig. 4.8)

**Fig. 4.8** Northern blot analysis of PRG-5 (11)
In situ hybridizations analysis highlighted tight regulation of mouse PRG-5 mRNA expression during brain development. PRG-5 was detected at embryonic day 17 (E17) and this expression is found in the hippocampal anlage, cortex and olfactory bulb. A strong hybridization signal was found in cortex and hippocampus at Post-natal day 0 and 5 (P0 and P5) (Fig. 4.9).

**Fig. 4.9** PRG-5 expression during embryonic and postnatal development. At embryonic day 17 (E17) hybridization signal for PRG-5 can be detected in the hippocampal anlage and olfactory bulb. At perinatal stages (E19 – postnatal day 10 (P10)), a strong hybridization signal is found in the cortex and hippocampus as well in dentate gyrus (dg). Scale bars 1.8 mm. CA, cornu ammonis; cx, cortex; Ob: olfactory bulb.

After that, studies using mRNA expression of PRG-5, normalized with HPRT or GAPDH housekeeping gene, revealed that expression of PRG-5 mRNA were significantly higher in brain than all other tissues, confirming that its expression is indeed brain-specific (Fig. 4.10).
Fig. 4.10 Multi-tissue real-time PCR analysis of PRG-5 mRNA shows the brain mainly expression of PRG-5 mRNA at P5. HPRT was used as reference gene. Error bars are standard deviations.

PRG-5 is expressed as early as embryonic day 14 (E14), peaks at postnatal day 0 (P0) (Fig. 4.11) and is principally expressed in neurons ad opposed to astrocytes and microglia of the brain hippocampus (Fig. 4.12)

Fig. 4.11 Quantification of PRG-5 mRNA in hippocampus mouse by real-time PCR. PRG-5 mRNA expression in the hippocampus of developing mouse brain, normalized to HPRT. Error bars are standard deviations.
Fig. 4.12 PRG-5 mRNA expression in neurons, astrocytes and microglia, normalized to GAPDH. Error bars are standard deviations.

4.3 Sub-cellular localization of PRG-5.

By PCR technique using total cDNA of mouse brain at P0, we generated a PRG-5 flag protein (DYKDDDD), as a tag fused at the C-terminus, instead of an EGFP fusion protein. Indirect immunofluorescence analysis of fixed cells revealed that the protein was localized to both intracellular membrane structures and to the plasma membrane (Fig. 4.13).

A

PRG-5 Flag

B

35 KDa

IB: anti-Flag

IB: anti-tubulin

Fig. 4.13 Overexpression of PRG-5 in HEK-293 cells. PRG-5 showed a punctuated distribution (white arrows) typically of the protein packed in vesicles. (B) Western blotting from total protein extraction from HEK-293. Asterisks indicate non-specific band.
In detail, intracellular localization of PRG-5 was analyzed by confocal microscopy, using Golgi and Rab9 vesicles marker (Fig. 4.14).

**Fig 4.14** PRG-5-Flag co-localizes with Golgi and Rab9 – vesicles marker in HEK-293 cells. (A and C) HEK-293 cells were transfected with pCDNA3-PRG-5-Flag. Fixed 24 hours later and investigated by confocal microscopy. (A) PRG-5 co-localize with Golgi marker (white arrows) and (B) partially with Rab-9 (white arrows). Scale bar are 10µm and 4µm respectively.
4.4 PRG-5 induces spine formation in primary neurons.

In light of the brain-specific expression in mouse, especially in the case of neurons, we performed a series of overexpression experiments in mouse hippocampal primary, cultivated between DIV 2 and 4. At this age, PRG-5 was widely expressed. The mRNA expression of PRG-5 was revealed by RT-PCR. I used a couple of primer to amplified PRG-5 and β-actin as control. In line 1 and 2, as template, I used cDNA from total mRNA of primary neurons at DIV 2 and 4 respectively, in line 3 H2O as negative control and in the line 4 a pcDNA-PRG-5 vector as positive control. The asterisk in line 4, respect the b-Actin primers, indicate an excess of primers and dNTPS used for the PCR (Fig. 4.15)

Fig. 4.15 RT-PCR from primary neurons at DIV 2 and 4 (line 1 and 2), H2O (line 3) and pcDNA-PRG-5 (line4) with primers to amplify PRG-5. M = Lambda marker digested by PstI.

Western-blot analysis of protein extracted from neurons at different days in vitro (DIV) with a PAP2D antibody, specific for PRG-5, revealed that PRG-5 is espressed at both DIV2 and DIV4. In particular, PRG-5 was detected in the membrane of the neurons, but not in their cytosolic fractions (Fig. 4.16 A). To study the overexpressed PRG-5 protein, I fused a Flag tag to the C-terminus of PRG-5. Membrane and cytosolic fractions from HEK-293 cells overexpressing PRG-5-Flag were tested using Western blot, and confirmed that both antibodies were able to detect PRG-5 specifically (Fig. 4.16 B and C).
Protein extraction was analyzed by Western blotting using PAP2D (PRG-5) and Flag antibodies. (A) Analyzed membrane (m) and cytosolic (c) fractions showed endogenous expression of PRG-5 in primary neurons, cultivated 2 or 4 days in vitro. (B and C) PRG-5-Flag overexpressed in HEK-293 cells was detected with Flag and PAP2D antibodies in the membrane fraction.

By means of confocal microscopy, a PRG-5-Flag signal was revealed on the body of the cell, in the proximal region of dendrites and at the distal neurites (Fig. 4.17). Higher magnification showed the presence of spine-like, mushroom-shaped and stubby structures along the plasma membrane. Distribution of PRG-5 along the plasma membrane appeared discontinuous, with a pronounced localization to many membrane protrusions (Fig. 4.17, white arrowhead).

The intracellular punctuated signal it was investigated in details using a α-clathrin vesicle, which is a specific vesicular carrier for membrane proteins. Clathrin spots were identified in the same area of PRG-5-Flag, suggesting a colocalization of PRG-5 with clathrin (Fig. 4.18, open arrows in central and right panel).
Fig. 4.17 Transfected primary neurons were fixed at DIV 2 and stained with antibodies to PRG-5-Flag (green) and MAP-2 (red). Scale bar represents 10 µm. Higher magnification of neuritis from a white box shows spine-like protrusions (white arrowheads). Scale bar represents 4 µm.

Fig. 4.18 PRG-5 (left panel) stained with antibodies to α-clathrin (central panel, open arrows). Merged panel shows PRG-5-induced spine structure (white arrowhead) and colocalization between PRG-5 and α-clathrin (open arrows). Scale bar represents 3 µm.
More in-depth analysis of PRG-5 was performed on primary neurons at DIV 2 or 4 by means of overexpression using markers for actin and tubulin. It was possible to distinguish swollen structures with a massive presence of spine structures containing PRG-5 (Fig. 4.19A and higher magnification, white arrowheads at the left panel). They originated from the plasma membrane, becoming more apparent along the processes. These protrusions were rich in polymerized actin and PRG-5-Flag was predominantly distributed uniformly along the thin neck and in the bulbous head of each single spine (Fig. 4.19A, higher magnification, central and merged panel), whereas tubulin structures appeared only along the processes of the neurites but was not in the bulbous head (Fig. 4.19B, white arrowheads). Primary neurons at DIV 4 overexpressing PRG-5-Flag showed the same distribution and shape of spines displayed in neurons at DIV 2 (Fig. 4.21 A). These types of spines were not found when EGFP alone was overexpressed in the same primary neurons (Fig 4.20 A and B and Fig. 4.21 B).
Fig. 4.20 (A and B) Overexpression of EGFP did not alter the morphology of the cells or induce spine formations along the neuritis. The scale bar represents 10 µm and 4 µm for higher magnifications.

Fig. 4.19 PRG-5 induces the dendritic spine formation during the first stages of neuronal differentiation. (A) Cultured primary neurons transfected with PRG-5-Flag (green) at DIV 2. Scale bar represents 10 µm. High-magnification insets show that PRG-5 (green) induces mature, mushroom-shaped spines along individual neurites that extend through the peripheral region, visualized by phalloidin (red). Arrowheads indicate spines with PRG-5 in the bulbous head. Scale bar represents 4 µm. (B) Higher magnification of primary neurons transfected with PRG-5-Flag (green) and visualized by tubulin (red). The white arrowheads indicate the heads of different spines induced by PRG-5. Scale bar represents 4 µm.
**Fig. 4.21** (A) Primary neurons transfected with PRG-5-Flag (green) at DIV 4 and visualized with actin (central bottom panels) or tubulin. Arrowheads indicate different mushroom-shaped and stubby spines. The scale bar represents 4 µm. (B) EGFP overexpressed in primary neurons at DIV 4 and detected using actin (central bottom panels) or tubulin. The scale bar represents 4 µm.
4.5 Analysis of residues within the C1 – C3 domains of PRG-5

Sigal Y. J. et al. have reported that single exchange of the amino acids Ser198 and His200 within the C2 domain and Arg246 within the C3 domain of PRG-3 in COS7 and HeLa cells is crucial for a correct functioning of the protein and resulted in an overproduction of the number of filopodia (58). However, despite the fact that analysis of the sequence homology between PRG-5 and PRG-3 showed a similarity higher than 85% (GeneBank), the residues within the C1-C3 domains, responsible for filopodia induction in PRG-3, were not completely conserved in PRG-5 (Tab. 1). In fact, although the amino acids Ser198 and Arg246 within the C2 and C3 domains, respectively, are both conserved in PRG-3 and PRG-5, in the case of His200 within the C2 domain of PRG-3, PRG-5 instead showed Glu195 (Tab. 1 and Fig. 4.22).

To understand if that residues, in PRG-5, have a role in the spine induction previously showed, I generated a series of proteins mutated at these single amino acids. The conservative amino acids Ser193 and Arg241 were exchanged with Trp193 and Glu241, respectively, while the non-conservative Glu195 was exchanged firstly with Gly195 and then with His195, to reproduce the same sequence of the PRG-3 C2 domain in PRG-5 (see also Tab. 1).

![Mouse PRG-5 structure. In red the three amino acids (Ser193, Glu195 and Arg241) exchanged. The double line represent the plasma membrane.](image.png)
Western blot analyses of mutated PRG-5-Flag construct overexpressed in cultured HEK-293 cells confirmed expression of all mutants at the predicted molecular mass of 35 KDa (Fig. 4.23).

![Western blot analysis of mutant proteins detected with a flag-specific antibody.](image)

**Fig. 4.23** Western blot analysis of mutant proteins detected with a flag-specific antibody.

Primary neurons were transfected with the same amount of PRG-5-flag wild type and PRG-5 Ser193Trp (PRG-5 S193W), -Glu195Gly (PRG-5 E195G), -Glu195His (PRG-5 E195H) and -Arg241Glu (PRG-5 R241E), fixed 24 hours later and investigated by confocal microscopy (Fig. 4.24). Primary neurons, overexpressing the PRG-5 protein with mutations at conservative amino acids S193W and R241E, as well non-conservative residue E195G, did not show any spines or membrane protrusions between DIV 2 and 4. Actin staining confirmed the absence of spines, filopodia or extracellular structures (Fig. 4.24 and 4.25).
Fig. 4.24 Neurons at DIV 2 overexpressed the S193W, E195G, R241E and E195H PRG-5 mutants and investigated by confocal microscopy. Merge colors showed PRG-5-Flag mutants and actin staining. Scale bar represents 10 µm. Higher magnification of neuritis from a white box. PRG-5-Flag antibody (green, central panels) and actin staining (red, right panels). Scale bar represents 4µm.
**Fig. 4.25** Primary neurons at DIV 4, transfected with mutants S193W, E195G and R241E were unable to induce spines respect the mutant E195H. Mutants are stained in green while red showed actin staining. Scale bar represents 4μm.
In detail, primary neurons transfected with mutants S193W, E195G and R241E displayed a dispersed fluorescence signal in the intracellular region, partially in the soma and along the neurites (Fig. 4.24 and 4.25). This dispersed fluorescence could be explained as an intracellular localization of the mutant proteins. Exchange of the amino acids Ser193, Glu195 and Arg241 may interfere with a correct protein folding, blocking the transport of the self-mutated protein to the plasma membrane. Nevertheless, the mutant E195H, as the PRG-5-Flag wild type, induced spine formation in neurons at DIV 2 and 4, respectively (Fig. 4.24 and 4.25, bottom panels).

The overexpression of PRG-5-Flag E195H in primary neurons showed a considerable number of spine-like structures along the soma and the neuronal processes. Each spine was rich in actin structure and a clear fluorescent signal from the induced mutant protein was distinguishable. In addition, sequence alignment of different PRG-5 species (GeneBank) revealed that Glu195 was highly conserved within the C2 domain, while all PRG family members displayed Ser193 and Arg241 as highly conserved within the C2 and C3 domains, respectively (Tab. 1).
5. Discussion

The members of PRG family define a subclass of the LPP superfamily, characterized by six membrane-spanning domains containing three extracellular loops within highly conserved enzymatic active domains (6). LPP-1 is able to degrade LPA, controlling the level of lipids in the extracellular space (10, 29, 30, 50) and expression pattern of LPP-1 is been found in different tissues, included in the brain (27, 31, 63).

Here, I have investigated the upstream region of PRG-1, identified as proximal promoter and successively, I characterized the last member of PRG family, plasticity-related gene-5 (PRG-5), able to induce spine-like structure in young primary neurons.

5.1 Proximal promoter of PRG-1

PRG-1 belongs to the family of plasticity related genes, which possesses homology to lipid phosphate phosphatases, and is part of the superfamily of lipid phosphate phosphatases/phosphotransferases (57). However, PRG-1 shows only a residual enzymatic activity for dephosphorylation of the superfamily specific substrates such as phosphatidate, lysophosphatidate or sphingosine-1-phosphate (10), which is due to amino acid exchanges in the catalytic motif that is shared by active PAP-2 family members (9). Principally, PRG-1 is involved in modulation of neuronal transmission via bioactive lipids, such as LPA acting via presynaptic LPA2 receptors, which are controlled by PRG-1 from the postsynaptic side (specifically at the glutamatergic junction). According to recent data, this action is mediated by PRG-1 by depleting the pool of bioactive lipids that act on the presynaptic LPA2 receptors (66).

The transcription of a eukaryotic gene is regulated by the combined action of multiple sequence-specific transcription factors, general transcription factors, histone modifiers, cofactors, and mediators that regulate transcription factor activity and chromatin structure. Our previous analysis of PRG-1 transcriptional regulation has shown that PRG-1 transcription is initiated at multiple transcription start sites. One sequence element 5'- at the translation start site (ATG) was detected. It mediates specific neuronal transcription in the human and rodent PRG-1 promoter (Geist et al., in progress). Interesting to note is that PRG-1 is transcribed via a type "null" core promoter neither TATA box nor initiator element, and no CCAAT box. Multiple start sites are
common for this promoter type (40). Computer aided analysis predicted also a 681 bp CpG island with 63.1% GC content 298 by upstream the first ATG of PRG-1. The existence of transcription start sites downstream of the first ATG of PRG-1 implies the existence of a second, alternative N-terminus. The shorter form of PRG-1 from second ATG needs to be confirmed, but resembles the homologous protein PRG-2 at the N-terminus (7), that is expressed prenatally and has a different expression pattern compared to PRG-1. Thus, the shorter form of PRG-1, lacking the N-terminal 49 amino acids, may substitutes the function of PRG-2 at postnatally stage. However, the function of N-terminal elongation remains to be clarified.

Recent study displayed the influence of Nex1, a homolog of the Drosophila proneural gene atonal belonging to the family of basic helix-loop-helix (bHLH) transcription factors, on PRG-1 expression. bHLH transcription factors bind to the consensus sequence CANNTG (E-Box), which has been identified in a variety of promoter and enhancer elements (35). Nex1 expression occurs at embryonic day 11.5 (55), and peaks in the first postnatal week parallel to dendritic arborisation and synaptogenesis of cortical neurons (2). Nex1 binds directly to at least on E-Box in the promoter region of PRG-1 and that it also mediates neurite outgrowth via PRG-1 expression in PC12 cells (72). Nex1 also regulates a molecular chaperone network through the expression of a set of heat shock proteins in the absence of stress, and connects neuronal differentiation with survival pathways (67). To determine the regulatory role of Nex1 in PRG-1 expression, I performed dual luciferase assays in primary cultured cells and subsequently analyzed elements displaying activation regarding Nex1 regulation.

In details, we identified a 450 bp minimal promoter conferring enhanced specific transcription in neurons. Computer aided analysis for identification of transcription factor binding sites delivered several candidates on the 450 bp promoter mediating the neuronal stimulation of transcription. A Nex1 independent regulation of PRG-1 which is mediated by a 450 bp promoter fragment, conferring enhanced specific transcription in neurons, was determined (Geist et al., in progress). This regulation can be demonstrated under organotypic conditions, which underlines that this sequence is sufficient for the peculiar neuronal expression of PRG-1 in neurons. Moreover, a Sp1 binding sites, sequence-specific transcription factor that binds GC and GT boxes to activate a wide range of viral and cellular genes, were found inside the core promoter of PRG-1. Sp1 is important both in transcription initiation and activation, and it can be regulated by multiple mechanisms in a cell type-specific and promoter context specific manner (5). Sp1 has been
linked to the maintenance of methylation-free CpG islands (34), and hypermethylation around Sp1 binding sites has been reported to reduce Sp1 binding, thereby decreasing transcription (75).

5.2 Plasticity-related gene 5 (PRG-5)

PRG-5 is an integral membrane protein containing 321 amino acids (aa) with both the N- and C-terminus present in the intracellular region that is representative of a class of five proteins with high homology to the lipid phosphate phosphatases (LPPs). Members of this class of proteins are characterized by highly conserved enzymatic active domains (6). LPP-1 is able to degrade LPA and control the level of lipids in the extracellular space (10, 29, 30, 50) and its expression pattern has been found in different tissues (31), including in the brain (27, 31, 63). Sequence alignment analysis revealed more than 95% similarity between human, rat and mouse PRG-5 and more than 80% homology with mPRG-3. Despite this, comparison of the relevant regions of PRG-3 and PRG-5 proteins with those of LPP family shows that crucial amino acids responsible for enzymatic ecto phosphatase activity are exchanged (26, 61, 64, 74). The His171 and His223 within the C2 and C3 consensus domains of LPP-1, essential for enzymatic function, were replaced in PRG-5 with Glu195 and Asn247 respectively (Tab. 1). Consequently, PRG-5 should not have any ectophosphatase activity, as recently shown for PRG-3 (51, 58). However, overexpression of PRG-5 induced spine-like structures in young primary neurons.

Mainly expressed in the brain (11), I confirmed that PRG-5 mRNA was presented as early as embryonic day 14 (E14) and peaked at postnatal day 0 (P0), while the PRG-5 protein was detected only in plasma membrane fraction of neurons at DIV 2 and 4.

In neurons at DIV 2, 24 hours after transfection, PRG-5-Flag protein was localized to both the intracellular region and plasma membrane. Immunostaining using a marker for α-clathrin vesicle showed a partial co-localization between PRG-5-Flag and α-clathrin, which suggests that PRG-5 may be recruited from these vesicles in the cytosol region to be transported into the plasma membrane. In the other hand, the plasma membrane showed the presence of many spine-like structures, probably PRG-5-Flag induced.

Dendritic spines observed in fixed brain tissue shows various shapes, and are generally classified into three types: the thin type having a slender neck and a small head, the mushroom type having
a short neck and a relatively large head, and the stubby type having no neck (Fig. 5.1). In living neurons, spine shapes easily interchange between the above three types. In other words, spine morphologies are snapshots of dynamic morphological changes. Therefore, not only the spine morphology but also its dynamic change should be elucidated to understand synaptic functions.

**Fig. 5.1** Schematic drawing of spine morphologies. Described in (73).

Hippocampal tissue prepared from P2-P7 animals and cultured 1-2 days in vitro generally show dendritic branches and lateral filopodia. When slices were cultured for 6-8 days in vitro, synaptogenesis and dendritic differentiation proceeded in vitro and spiny protrusions with complex shapes were clearly present (15). Dendrites of immature neurons, cultured for DIV 7 showed thin and long protrusions, clearly filopodia while neurons cultivated for DIV 21 were mature and covered with spines (54). On the other hand, dendrites of hippocampal neurons at DIV 10, transfected with EGFP, were covered with thin, long filopodia-like protrusions (44).

However, in our study, overexpression of PRG-5-Flag led to a large number of spines in neurons as young as DIV 2 and DIV 4. The spines were found with a mushroom and stubby shape, and the bulbous head contained PRG-5-Flag and polymerized actin. Filament of tubulin was detected only around the soma and along the neuritis, as expected. These types of membrane protrusions were not labelled when EGFP alone was expressed in these cells and suggest that a pathway for spine induction may exist that depend on PRG-5 expression. Stable spines could emerge on developing dendrites from pre-existing spiny structures or from dendrite shafts by *de novo* extension (15). In addition, newly formed spiny-like protrusions, developed over 48 hours, are
morphologically indistinguishable from preexisting spines, complicating the correct identification of the provenience of the mushroom spines already observed (76). Although our own observations are of primary neurons and not fixed tissue, it is interesting to note that neurons from E18 mice, cultivated at least 1 day in vitro, were able to produce mushroom and stubby spines after 24 h of overexpression of PRG-5, mainly along the neurites of the cell. This was still observable after 4 days in vitro, but not when the neurons were transfected with an empty EGFP as control.

Although the mechanism by which PRG-5 regulates the formation of spines requires further investigation, mutational studies provide insight into the possible mechanisms responsible. Unlike an earlier study, which analyzed a truncated PRG-5 protein, in which the C-terminus, after deletion, did not allow for filopodia induction in N1E-115 cells (11), we investigated the “catalytic sites” of PRG-5 using single-point mutations, as previously reported for PRG-3. In details, PRG-3 overexpressed in COS7 and HeLa cells is able to increase the number of filopodia and the amino acids Ser193, His200 and Arg246, within the catalytic domains C1 – C3, were responsible for this function (58). However, despite the fact that PRG-5 displays high sequence homology with PRG-3, it lacks critical residues present in the latter. In particular, instead of His200 within the C2 domain of PRG-3, PRG-5 instead shows Glu195 (Tab. 1).

Using direct mutagenesis, we substituted conservative Ser193 (C2 domain) and Arg241 (C3 domain) with Trp193 and Glu241 respectively and the non-conservative Glu195 (C2 domain), firstly in Gly195 and then in His195, in PRG-5 to replicate the C2 domain of PRG-3 (Tab. 1). The mutants S193W, R241E and E195G failed to induce spine formation in primary neurons at DIV 2 (Fig. 4.22) and 4 (Fig. 4.23). The predominant intracellular fluorescence distribution detected in the primary neurons could be the consequence of an incorrect folding of the mutated proteins, which may have hindered the transport of mutated PRG-5 to the plasma membrane. PRG-5 with the Glu195 mutated in His195, to reproduce the same C2 domain of PRG-3, was again able to induce spine formation in primary neurons.
This diagram illustrates the essential features of the three models of spinogenesis. In the Sotelo model (a), spines emerge independently of the axonal terminal. In the Miller/Peters model (b), the terminal actually induces the formation of the spine. Finally, in the filopodial model (c), a dendritic filopodium captures an axonal terminal and becomes a spine. (73).

Although the mechanisms involved in spine regulation remain unclear, three models for spinogenesis have been proposed, known as the Sotelo, Miller/Peters and Filopodia models (73) (Fig. 5.2). The Miller/Peters and Filopodia models suggest that a physical axon-dendrite interaction is required for the formation of spines (73), whereas the Sotelo model describes spines that emerge from dendrites independently of the axon. First, synapses are made on the dendritic shafts, and immature spines can be recognized by their flocculent material. Most of these spines are ‘stubbies’. In the second stage, the presynaptic region of the axon shows a swelling as synaptic vesicles accumulate. In the third stage, many spines are thin or mushroomshaped, with a lollipop shape and a clear neck, and axonal terminals have well developed varicosities (Fig. 5.2 B and C).

This hypothesis is different from the Sotelo model (Fig. 5.2 A), which proposes that the terminal has a minor role in spinogenesis. It is possible that Purkinje cells and pyramidal neurons use completely different spinogenesis strategies, as they are very different cell types with seemingly
different circuit functions (for example, excitatory versus inhibitory). At the same time, although there is consensus regarding the time course of events in pyramidal neurons, we feel that the available data on pyramidal spinogenesis are circumstantial and that the Miller/Peters hypothesis needs to be properly demonstrated (73).

PRG-5-Flag induces spine formation in neurites of immature primary neurons independently of an axon-dendrite interaction. Furthermore, our findings suggest that a necessary condition for PRG-5 to induce spines in primary neurons is a correct positioning of the protein at the plasma membrane. Given this, PRG-5, as a multispansng membrane protein, can induce a membrane deformation and consequently spine induction. A possible curvature of membrane was proposed for the I-BAR proteins (28). In the latter case, filopodia initiation could occur through membrane-deforming proteins that contain I-BAR (inverse Bin/Amphiphysin/Rvs) domains. Electrostatic interactions between the positively charged poles of I-BAR domains and the negatively charged PI(4,5)P2 head groups induce clustering of PI(4,5)P2 and generation of membrane curvature due to the convex geometry of the lipid-binding interface of the domain (28). Specifically in neurons, depletion of I-BAR on the protein IRSp53 affect dendritic spine morphogenesis (14). This is consistent with the observation that primary neurons transfected with PRG-5 mutated protein exhibit no spines, and that proteins were detected in the cytosol and partially in the membrane.

![PRG-5 Flag Tubulin Merge](image)

**Fig. 5.3** (A) HEK-293 cells transfected with pCDNA3-PRG-5-Flag (green) and analyzed by confocal microscopy. Tubulin structure is visualized in red. (B) Higher magnification of boxed area. Scale bar represents 10 and 4µm respectively.
Moreover, a transient overexpression of PRG-5 in HEK-293 cells, resulted in a pronounced morphological changes of the plasma membrane. PRG-5-Flag protein appeared packaged inside capsules that are all around the cells, as an incontrollable exocytosis (Fig. 5.3), may due to an extreme rearrangement of the plasma membrane.

In conclusion, we have found that mouse PRG-5, an integral transmembrane protein, is brain-specific and able to induce mushroom and stubby spines in immature primary neurons prior to the start of the physiological process of spinogenesis. Although we do not know the basis of this spine induction, our results suggest PRG-5 plays a critical role in spine induction, and that a basic prerequisite for this functioning is correct positioning at the plasma membrane.
5. Bibliography


7. Publication in progress and National congress

Publication in progress

**Coiro P.,** Velmans T., and Bräuer A.U.

Plasticity-related gene 5 induces spine formation in immature primary neurons. Submitted on October 2010 to MCB.

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Determination of PRG-1: Transcriptional Regulation Independent from Nex1/Math2 Mediated Activation. Submitted on September 2010 to Journal of Neuroscience.

National congress

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Genomic organization of PRGs in humans: functional evidence for the promoter region of PRG-1. **Poster**

**P. Coiro,** A.U. Bräuer, N.E. Savaskan, R. Nitsch
June 2008, BNF Berlin
PRG-5 in cultured neurons inducts dendritic-spines formation. *Poster*

**P. Coiro, T. Velmans, A.U. Bräuer**

June 2010, BNF Berlin

PRG-5 induces spine in immature primary neurons. *Poster*
8. Declaration

I declare that this thesis is my original and unaided intellectual work. It is being submitted for the award of the degree of Doctor of Philosophy (PhD) at the Freie Universität of Berlin. It has not been submitted before for any degree or examination in any other university here in Germany or anywhere else in the world.

Berlin, den………………………………03.01.2011

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Pierluca Coiro
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