

BMC Anesthesiology

Open Access

Notch signaling activation is critical to the development of neuropathic pain

Keliang Xie^{1*†}, Feng Qiao^{2*†}, Yanyan Sun³, Guolin Wang¹ and Lichao Hou^{3*}

Abstract

Background: Nerve injury-induced neuropathic pain is a major health problem worldwide. Notch signaling is a highly conserved pathway in evolution, which has an important role in synaptic plasticity and inflammation in central nervous system. The present study was designed to investigate the potential role of notch signaling in the development of neuropathic pain.

Methods: The neuropathic pain was induced by spared nerve injury (SNI) in rats. The activation of notch signaling in the lumbar spinal dorsal horn was measured. DAPT, an inhibitor of notch signaling, was intrathecally (i.t.) administered before SNI or after appearance of pain sensitivity. Moreover, Jagged-1 (JAG-1) peptide, a ligand of notch signaling, was i.t. administered to normal rats. The mechanical allodynia was assessed by von Frey test.

Results: Here, we found that DAPT administered 0.5 h before SNI operation could significantly prevent the decrease of mechanical paw withdrawal threshold (PWT) for more than 4 weeks (P < 0.05 vs. SNI group). DAPT administered after appearance of pain sensitivity could also significantly reverse the decrease of mechanical PWT in a dose-dependent manner (P < 0.05). In addition, administration of Jagged-1 (JAG-1) peptide significantly decreased the mechanical PWT of normal rats in a dose-dependent manner (P < 0.05).

Conclusions: Therefore, notch signaling activation might contribute to the development of neuropathic pain. This study might provide a new therapeutic target for neuropathic pain.

Keywords: Neuropathic pain, Notch signaling pathway, Mechanical allodynia

Background

Neuropathic pain is a chronic pain resulting from dysfunction or damage of the nerve fibers of peripheral or central nervous system (CNS) [1]. The mechanism is complicated and unclear, which is associated with hyperexcitability in the affected dorsal root ganglion (DRG) neurons, atrophic changes and a switch in neurotransmitter phenotype in the central afferent terminal, aberrant myelination, alterations in synaptic plasticity and excitatory and inhibitory mechanisms in spinal cord, and loss of inhibitory interneurons and modifications of

* Correspondence: xiekeliang2009@hotmail.com; 58159529@qq.com; houlc9534@hotmail.com

brain input to spinal cord [1-7]. Although there is significant improvement in treatment, neuropathic pain frequently remains unresponsive to all treatment modalities [8]. Notch signaling is a highly conserved pathway in evolution, which often regulates cell-fate decisions in developing nervous system and has an important role in synaptic plasticity in adult CNS [9-12]. Numerous studies have demonstrated that notch signaling is crucial for many biological processes such as development, immunology, inflammation, tumor formation and memory [11,13-17]. Recent studies show that notch signaling activation might contribute to neuronal death, inhibition of neurite growth, more dendritic branching, generation and activation of microglial cells and astrocytes, differentiation of oligodendrocyte progenitors and demyelination in both peripheral nervous system and CNS [9,18-24]. Moreover, notch signaling controls the choice between excitatory and inhibitory cell fates in developing spinal cord, and its activation can promote the generation of



© 2015 Xie et al.; licensee BioMed Central. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly credited. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

[†]Equal contributors

¹Department of Anesthesiology, Tianjin Institute of Anesthesiology, General Hospital of Tianjin Medical University, Tianjin 300052, China

²Department of Orthopedics of Integrated Traditional Chinese and Western Medicine, Hong Hui Hospital, Xi'an Jiaotong University College of Medicine, Xi'an 710054, China

³Department of Anesthesiology, Xijing Hospital, Fourth Military Medical University, Xi'an 710032, Shaanxi Province, China

excitatory neurons from the sensory interneuron progenitors [25]. In addition, the expression and activity of notch signaling are significantly increased after nerve injury [26]. Ligands such as Delta and Jagged bind to Notch receptors, resulting in proteolytic cleavage of Notch into two sections: an extracellular domain and a transmembrane domain. The latter cleavage is completed by the γ -secretase enzyme and ADAM, which results in the release of a Notch intracellular domain (NICD).

Therefore, we hypothesized that notch signaling activation might contribute to the development of neuropathic pain. In the present study, we firstly investigated the activation of notch signaling in the lumbar spinal dorsal horn. Then, the effects of a notch signaling inhibitor DAPT administered at different times on the mechanical allodynia were measured in a rat model of spared nerve injury (SNI)-induced neuropathic pain. Last, we studied the effects of a notch signaling activator Jagged-1 (JAG-1) peptide on the mechanical allodynia in normal rats. This study may provide a new therapeutic target for neuropathic pain.

Methods

Animals

All experiments were performed on adult male Sprague-Dawley rats weighing 200-250 g (Laboratory Animal Center of the Tianjin Medical University, Xi'an, China). The animals were housed in plastic boxes at 22-26°C with food and water available *ad libitum*. A 12:12 h light/dark cycle with lights on at 8:00 was maintained and testing was done between 9:00 and 18:00. Prior to experimental manipulation, animals were allowed to acclimate to the housing facilities and were handled daily at least for 3 d. All experimental protocols and animal handling procedures were approved by the Tianjin Medical University Animal Care and Use Committee and were in accordance with the guidelines for the ethical treatment of animals established by the International Association for the Study of Pain. All efforts were made to minimize animal suffering and to reduce the number of animals used.

Intrathecal catheterization and drug delivery

A permanent intrathecal (i.t.) catheter (PE-10 polyethylene tube, BD Biosciences, Franklin Lakes, NJ) was inserted through the gap between $T_3 \sim_4$ vertebrae and extended slowly to the subarachniod space of lumbar enlargement (L₄ and L₅ segments) under pentobarbital sodium anesthesia [40 mg/kg, intraperitoneal (i.p.)] [27]. The catheter was filled with sterile saline (approximately 4 μ L), and the outer end was plugged. All animals appeared to be free of infection through gross inspection. The cannulated animals were allowed to recover for 4 days. Motoric integrity was assessed in all animals using the righting reflex and

inclined plane test. Animals showing any neurological deficits were excluded from the following experiment.

Drugs were injected over a period of 1 min via the catheter at a volume of 10 μ L, followed by 5 μ L sterile saline for flushing. DAPT (Sigma-Aldrich, St. Louis, MO, USA) and JAG-1 peptide (CDDYYYGFGCNKFC RPR) (AnaSpec, San Jose, CA, USA), as a potent inhibitor and activator of notch signaling, respectively, were both freshly dissolved in dimethyl sulfoxide (DMSO) at a concentration of DAPT (25, 50 or 100 μ M) and at a concentration of JAG-1 peptide (1, 10 or 100 μ M) [19]. The DMSO or scrambled peptide (SC)-JAG-1 (RCGPD CFDNYGRYKYCF) (AnaSpec, San Jose, CA, USA) was used as a control. The location of the distal end of this catheter was verified at the end of each experiment by injection of pontamine sky blue via the catheter.

Spared nerve injury (SNI) model

The neuropathic pain was induced by left SNI model as previously described [28]. Briefly, under pentobarbital sodium anesthesia, an incision was made on the lateral thigh and the underlying muscle was separated to expose sciatic nerve. The three terminal branches of sciatic nerve (tibial, common peroneal and sural nerves) were carefully separated. After separation, the tibial and common peroneal nerves were tightly ligated with 5.0 silk, and then 2–3 mm of the nerves distal to the ligation was removed. The muscle and skin incisions were then closed separately. Sham operation was performed identically without the passage of ligatures or transection of the nerves.

Assessment of notch signaling activity

The activation of notch signaling was measured by western-blot of NICD. The lumbar spinal dorsal horn tissues were harvested, sonicated/homogenized and centrifuged. The protein was extracted from the above preparations, and loaded and separated on a 10% (w/v) SDS-polyacrylamide gel. After transfer, blots were incubated overnight at 4°C with separately with NICD (1:1000; Abcam, London, UK) and beta-actin (1:10000; Chemicon, Temecula, CA, USA) primary antibodies. The membrane was washed and primary antibodies were detected with secondary antibody conjugated to horseradish peroxidase for 1 h at room temperature. Blots were visualized using enhanced chemiluminescence (ECL; Roche, Basel, Switzerland).

Assessment of mechanical allodynia

The mechanical allodynia was assessed by von Frey test as previous described [29]. Mechanical allodynia of rats was determined by paw withdrawal threshold (PWT) in response to mechanical stimuli produced by using a calibrated series of von Frey filaments (Stoelting, Chicago, IL, USA). The animals were adapted to the testing situation for at least 30 min before stimulation was initiated. During the test, the rats were placed on a metal mesh floor covered with the same plastic box and von Frey filaments were applied from the underneath of metal mesh floor to the lateral plantar surface of the paw (the area innervated by sural nerve). Each filament was presented perpendicularly against the paw, with sufficient force to cause slight bending, and held 2-3 s. The filament was applied only when the rat was stationary and standing on all four paws. A withdrawal response was considered valid only if the hindpaw was completely removed from the customized platform. Lifting of the paw due to normal locomotor behavior was ignored. The monofilaments were applied with increasing force until the rat withdrew the paw. Each hair was applied 10 times at 5 s intervals. The bending force value of the von Frey filament that caused a 50% occurrence of paw withdrawal reflex by 10 times stimuli was expressed as the PWT. After the threshold was determined for the left hindpaw, the same testing procedure was repeated on the other hindpaw at 5 min interval. Second and third testing trials were run for both hindpaws, respectively. If the withdrawal threshold in the second or third trial did not match the withdrawal threshold of the previous testing trial in a given hindpaw, the next large hair in the series was tested. This was done until the withdrawal threshold in the three successive trials matched. To avoid inter-experimenter differences and subjective bias, all the behavioral observations were performed by one who was blind to the treatment.

Statistical analysis

Data are expressed as mean \pm standard deviation (SD). Statistical comparisons between groups were done using univariate ANOVA or ANOVA for repeated measurements. The statistical analysis was performed with *SPSS* 16.0 software (SPSS Inc, Chicago, IL). A *P* value of less than 0.05 was considered statistically significant for all tests.

Results

Notch signaling is activated after SNI operation

In this study, we firstly investigated the notch signaling activity at different time points after SNI operation. This present study showed that NICD expression in the dorsal horn of lumbar spinal cord was markedly increased until 28 days following peripheral nerve injury (Figure 1). This result suggests that notch signaling is activated during the development of neuropathic pain.

Administration of DAPT before SNI operation significantly prevents the development of mechanical allodynia in neuropathic pain

In the present study, we then investigated the effects of notch signaling pathway inhibitor DAPT pretreatment



on mechanical allodynia in a rat model of SNI-induced neuropathic pain. In this experiment, the mechanical allodynia in all animals was measured at 24 h before (baseline) as well as 7 d, 14 d, 21d and 28 d after SNI or sham operation. The SNI-challenged animals developed a marked hypersensitivity to innocuous mechanical stimulation of the lateral surface of the hindpaw (sural nerve skin area) (Figure 2). The hindpaw contralateral to the operation was tested over the whole period and did not demonstrate any statistically significant change in mechanical PWT from pre-operation baseline (data not shown). The mechanical PWT from 7 d to 28 d after SNI operation decreased significantly (Figure 2, P < 0.05vs. Sham group, n = 6 per group). A single administration of DAPT (50 µM, 10 µL, i.t.) 0.5 h before SNI operation significantly improved the mechanical PWT for more than 28 d after the operation (Figure 2, P < 0.05 vs. SNI group, n = 6 per group). In addition, we found that



10 μ L, i.t.) was 0.5 h before SNI operation. The mechanical mechanical paw withdrawal threshold (PWT) was measured at 24 h before (baseline) as well as 7 d, 14 d, 21 d and 28 d after SNI or Sham operation. The data represent mean \pm SD (n = 6 per group). *P < 0.05 versus Sham group; †P < 0.05 versus SNI group. SNI: spared nerve injury; BL: baseline; g: gram; d: day.

administration of DAPT had no effect on the mechanical PWT of normal rats (*data not shown*). This result suggests that early inhibition of notch signaling can prevent the development of mechanical allodynia in neuropathic pain.

Administration of DAPT after appearance of pain sensitivity significantly reverses the mechanical allodynia in neuropathic pain

We then investigated the effects of notch signaling pathway inhibitor DAPT posttreatment on mechanical allodynia in SNI-induced neuropathic pain. In the preliminary experiment, we found that the animals developed significant mechanical allodynia 3 d after SNI operation (data not shown). A single administration of DAPT (50 µM, 10 µL, i. t.) after appearance of pain sensitivity (80 h after SNI operation) significantly increased the mechanical PWT from 3 h to 72 h after DAPT administration (Figure 3A, P < 0.05vs. SNI group, n = 6 per group). Moreover, different concentrations of DAPT (25, 50 and 100 µM) were administered after the appearance of pain sensitivity, and the mechanical PWT was evaluated 24 h after DAPT administration. As shown in Figure 3B, administration of DAPT dose-dependently increased the mechanical PWT of neuropathic pain rats (P < 0.05). These results suggest that late inhibition of notch signaling pathway can also reverse the mechanical allodynia of neuropathic pain in a dosedependent manner.

Administration of Jagged-1 peptide, a ligand of notch signaling pathway, can induce neuropathic pain-like behavior in normal rats

To further investigate the role of notch signaling in neuropathic pain, the JAG-1 peptide, a ligand of notch signaling pathway, were i.t. administered of 1 μ M, 10 μ M, 100 μ M to normal animals, and the mechanical PWT was measured 3 h, 6 h, 12 h, 24 h, 36 h, and 48 h after JAG-1 administration. The SC-JAG-1 peptide was administered as a negative control. As shown in Figure 4, a single administration of JAG-1 peptide dose-dependently decreased the mechanical PWT of normal animals from 3 h to 36 h when compared with SC-JAG-1 peptide group (P < 0.05, n = 6 per group). The decreased PWT after the injection of JAG-1 peptide was restored 48 h later. The results further suggest that activation of notch signaling contributes to the development of neuropathic pain.

Discussion

In the present study, we firstly found that notch signaling in dorsal horn of lumbar spinal cord was activated during the development of neuropathic pain. Early inhibition of notch signaling can prevent the development of SNI-induced neuropathic pain. Late inhibition of notch signaling can reverse the mechanical allodynia of



neuropathic pain. Furthermore, activation of notch signaling can induce the mechanical allodynia of normal animals. Therefore, notch signaling activation might be critical to the development of neuropathic pain.

Injury in nervous system may result in chronic neuropathic pain characterized by increased sensitivity to painful stimuli (hyperalgesia), the perception of innocuous stimuli as painful (allodynia) and spontaneous pain [2,8]. It is well known that a characteristic symptom of neuropathic pain is mechanical allodynia [8]. The SNI model has proved to be robust, with substantial and prolonged changes in mechanical sensitivity that closely mimic many features of clinical neuropathic pain [28,29]. The model may provide a useful tool for identifying the underlying mechanisms involved in the development of neuropathic pain and as an additional screen





for the efficacy of new treatments. In this study, we found that all the animals developed significant mechanical allodynia in SNI-induced neuropathic pain, which is consistent with other studies [28,29]. Many potential mechanisms have been studied which may contribute to the pathogenesis of neuropathic pain. It has been reported that multiple mechanisms at multiple sites may operate either alone or together or at different time courses to produce the complex clinical characteristics [8]. These include changes in terminal and peripheral sensitization, phenotypic switches and excitability of injured axons, hyperexcitability in the affected DRG neurons, aberrant myelination (splitting, detachment and loss of myelin), synaptic plasticity in spinal cord, loss of inhibitory interneurons and modifications of brain stem input to spinal cord, and so on [1-7]. Alterations in the balance of number and/or transmission properties of these excitatory and inhibitory interneurons are thought to be major contributing factors for chronic sensory neuropathies such as hyperalgesia and allodynia [30,31]. In addition, sensory information from the periphery is integrated and transduced by excitatory and inhibitory interneurons in the spinal dorsal horn. However, the key mechanisms that control the induction and maintenance of neuropathic pain remain unclear.

Notch is a cell-surface receptor that regulates cell-fate decisions in developing nervous system and has important roles in synaptic plasticity in adult CNS [9-11]. Binding of ligands such as Delta and Jagged results in proteolytic cleavages of notch: first in an extracellular domain and then in a transmembrane domain [13,14]. The latter cleavage is accomplished by the γ -secretase enzyme complex, resulting in the release of a notch intracellular domain (NICD) that translocates into the

nucleus, where it regulates transcription [13,14]. All of the different components necessary for notch signaling, such as ligands, receptors, and enzymes involved in notch receptor cleavage, are expressed in the adult CNS as well as increase significantly after nerve injury and participate in its reparation [11,26]. Notch signaling is a highly conserved pathway in evolution, which is crucial for many biological processes such as development, immunology, inflammation, vasculogenesis, tumor formation, and learning and memory [11,13-17]. Recent findings suggest notch signaling activation can contribute to generation and activation of microglial cells and astrocytes, inhibition of neurite growth, more dendritic branching, differentiation of oligodendrocyte progenitors and demyelination in both peripheral nervous system and CNS [9,18-24]. Moreover, notch signaling pathway controls the choice between excitatory and inhibitory cell fates in the developing spinal cord, and its activation can promote the generation of excitatory neurons from the sensory interneuron progenitors [25]. To investigate the roles of notch signaling in neuropathic pain, a gamma-secretase enzyme (a key enzyme of notch signaling pathway) inhibitor DAPT was i.t. administered before or after appearance of pain sensitivity in the rat model of SNI-induced neuropathic pain. The results showed that early or late inhibition of notch signaling could prevent or reverse the mechanical allodynia of rats with neuropathic pain in a dose-dependent manner. In addition, administration of JAG-1 peptide (a ligand of notch signaling pathway) could induce the mechanical allodynia in normal rats in a dose-dependent manner. These results suggest that notch signaling activation contributes to the development of neuropathic pain.

The notch family includes 4 receptors with different functions, as indicated by the phenotypes of various notch transgenic mouse lines. These 4 receptors are expressed unequally in various cell types. Notch receptors and ligands have variable expression in various cell types. The notch signaling in the sciatic nerve, spinal dorsal horn and DRG neurons is markedly increased following peripheral nerve injury. Therefore, the notch signaling in neurons should have an important role in the development of neuropathic pain. DAPT is a nonspecific inhibitor of notch receptors. Therefore the study does not specify which notch receptor(s) are involved in allodynia. Moreover, Jagged 1 is also a non-specific ligand of Notch receptors. Moreover, Jagged 1 may have different affinities for different notch receptors, depending on cellular context and post-translational modifications. Also, the extracellular Jag1 peptide will activate notch signaling indiscriminately in neurons (both inhibitory and excitatory) and glia. However, one must keep in mind the widespread and varied expression of notch receptors and ligands. What will be the effect of a systemic

inhibition of notch signaling on other events where notch plays a key role in stem cell renewal? For example, hippocampus neurogenesis, that plays a key role in memory, will be also inhibited by notch inhibition.

Limitations of this study

The notch family includes 4 receptors with different functions. DAPT is a non-specific inhibitor of notch receptors. Therefore the study does not specify which notch receptor(s) are involved in allodynia. Moreover, Jagged 1 is also a non-specific ligand of notch receptors. Moreover, Jagged 1 may have different affinities for different notch receptors. Also, the extracellular Jagged 1 peptide will activate notch signaling indiscriminately in neurons (both inhibitory and excitatory) and glia. Moreover, the roles of notch signaling pathway in thermal or cold allodynia of neuropathic pain are still further studied. In addition, its roles in inflammatory pain are unclear. The underlying mechanisms are also the future research direction.

Conclusion

In conclusion, notch signaling activation might contribute to the development of neuropathic pain. This study might provide a new therapeutic target for neuropathic pain.

Key message

- 1. Notch signaling in spinal dorsal horn is activated in neuropathic pain.
- 2. Early inhibition of notch signaling pathway prevents the mechanical allodynia in neuropathic pain.
- 3. Late inhibition of notch signaling pathway reverses the mechanical allodynia in neuropathic pain.
- 4. Notch signaling activation induces the mechanical allodynia in normal animals.

Abbreviations

CNS: Central nervous system; DMSO: Dimethyl sulfoxide; DRG: Dorsal root ganglion; i.t.: Intrathecal; JAG-1: Jagged-1; NICD: Notch intracellular domain; PWT: Paw withdrawal threshold; SC-JAG-1: Scrambled Jagged-1; SNI: Spared nerve injury.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

FQ, YS performed the experiment and summarized the data, and wrote the manuscript and edited the manuscript. KX designed and performed the experiment, summarized and analyzed the data and edited the manuscript. GW analyzed the data and edited the manuscript. LH designed the experiment, analyzed the data and edited the manuscript. All authors read and approved the final manuscript.

Acknowledgements

This work was supported by research grants from the National Natural Science Foundation of China (81101409 and 81471842), the Natural Science Foundation of Tianjin, China (13JCQNJC11400), the Foundation of Tianjin Bureau of Public Health (2011KZ108), and the Xi'an City Social Development Project (SF1026(1)).

Received: 1 April 2014 Accepted: 10 March 2015 Published online: 28 March 2015

References

- Scholz J, Woolf CJ. The neuropathic pain triad: neurons, immune cells and glia. Nat Neurosci. 2007;10(11):1361–8.
- Zimmermann M. Pathobiology of neuropathic pain. Eur J Pharmacol. 2001;429(1–3):23–37.
- Willis WD. Role of neurotransmitters in sensitization of pain responses. Ann N Y Acad Sci. 2001;933:142–56.
- Woolf CJ. Evidence for a central component of post-injury pain hypersensitivity. Nature. 1983;306(5944):686–8.
- Moore KA, Kohno T, Karchewski LA, Scholz J, Baba H, Woolf CJ. Partial peripheral nerve injury promotes a selective loss of GABAergic inhibition in the superficial dorsal horn of the spinal cord. J Neurosci. 2002;22(15):6724–31.
- Porreca F, Ossipov MH, Gebhart GF. Chronic pain and medullary descending facilitation. Trends Neurosci. 2002;25(6):319–25.
- Gillespie CS, Sherman DL, Fleetwood-Walker SM, Cottrell DF, Tait S, Garry EM, et al. Peripheral demyelination and neuropathic pain behavior in periaxin-deficient mice. Neuron. 2000;26(2):523–31.
- Baron R, Binder A, Wasner G. Neuropathic pain: diagnosis, pathophysiological mechanisms, and treatment. Lancet Neurol. 2010;9(8):807–19.
- Louvi A, Artavanis-Tsakonas S. Notch signalling in vertebrate neural development. Nat Rev Neurosci. 2006;7(2):93–102.
- Latasa MJ, Cisneros E, Frade JM. Cell cycle control of Notch signaling and the functional regionalization of the neuroepithelium during vertebrate neurogenesis. Int J Dev Biol. 2009;53(7):895–908.
- 11. Costa RM, Drew C, Silva AJ. Notch to remember. Trends Neurosci. 2005;28(8):429–35.
- 12. Rusanescu G, Mao J. Notch3 is necessary for neuronal differentiation and maturation in the adult spinal cord. J Cell Mol Med. 2014;18(10):2103–16.
- Kopan R, Ilagan MX. The canonical Notch signaling pathway: unfolding the activation mechanism. Cell. 2009;137(2):216–33.
- 14. Fortini ME. Notch signaling: the core pathway and its posttranslational regulation. Dev Cell. 2009;16(5):633–47.
- Woodhoo A, Alonso MB, Droggiti A, Turmaine M, D'Antonio M, Parkinson DB, et al. Notch controls embryonic Schwann cell differentiation, postnatal myelination and adult plasticity. Nat Neurosci. 2009;12(7):839–47.
- Gridley T. Notch signaling in vascular development and physiology. Development. 2007;134(15):2709–18.
- 17. Shih le M, Wang TL. Notch signaling, gamma-secretase inhibitors, and cancer therapy. Cancer Res. 2007;67(5):1879–82.
- Arumugam TV, Cheng YL, Choi Y, Choi YH, Yang S, Yun YK, et al. Evidence that gamma-secretase-mediated Notch signaling induces neuronal cell death via the nuclear factor-kappaB-Bcl-2-interacting mediator of cell death pathway in ischemic stroke. Mol Pharmacol. 2011;80(1):23–31.
- Arumugam TV, Chan SL, Jo DG, Yilmaz G, Tang SC, Cheng A, et al. Gamma secretase-mediated Notch signaling worsens brain damage and functional outcome in ischemic stroke. Nat Med. 2006;12(6):621–3.
- Grandbarbe L, Michelucci A, Heurtaux T, Hemmer K, Morga E, Heuschling P. Notch signaling modulates the activation of microglial cells. Glia. 2007;55(15):1519–30.
- Yuan TM, Yu HM. Notch signaling: key role in intrauterine infection/ inflammation, embryonic development, and white matter damage? J Neurosci Res. 2010;88(3):461–8.
- Sestan N, Artavanis-Tsakonas S, Rakic P. Contact-dependent inhibition of cortical neurite growth mediated by notch signaling. Science. 1999;286(5440):741–6.
- Jurynczyk M, Jurewicz A, Bielecki B, Raine CS, Selmaj K. Overcoming failure to repair demyelination in EAE: gamma-secretase inhibition of Notch signaling. J Neurol Sci. 2008;265(1–2):5–11.
- Morrison SJ, Perez SE, Qiao Z, Verdi JM, Hicks C, Weinmaster G, et al. Transient Notch activation initiates an irreversible switch from neurogenesis to gliogenesis by neural crest stem cells. Cell. 2000;101(5):499–510.
- Mizuguchi R, Kriks S, Cordes R, Gossler A, Ma Q, Goulding M. Ascl1 and Gsh1/2 control inhibitory and excitatory cell fate in spinal sensory interneurons. Nat Neurosci. 2006;9(6):770–8.
- Givogri MI, de Planell M, Galbiati F, Superchi D, Gritti A, Vescovi A, et al. Notch signaling in astrocytes and neuroblasts of the adult subventricular zone in health and after cortical injury. Dev Neurosci. 2006;28(1–2):81–91.

- Yan LH, Hou JF, Liu MG, Li MM, Cui XY, Lu ZM, et al. Imbalance between excitatory and inhibitory amino acids at spinal level is associated with maintenance of persistent pain-related behaviors. Pharmacol Res. 2009;59(5):290–9.
- 28. Decosterd I, Woolf CJ. Spared nerve injury: an animal model of persistent peripheral neuropathic pain. Pain. 2000;87(2):149–58.
- Tegeder I, Costigan M, Griffin RS, Abele A, Belfer I, Schmidt H, et al. GTP cyclohydrolase and tetrahydrobiopterin regulate pain sensitivity and persistence. Nat Med. 2006;12(11):1269–77.
- Woolf CJ, Shortland P, Coggeshall RE. Peripheral nerve injury triggers central sprouting of myelinated afferents. Nature. 1992;355(6355):75–8.
- Fitzgerald M. The development of nociceptive circuits. Nat Rev Neurosci. 2005;6(7):507–20.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar

BioMed Central

(

• Research which is freely available for redistribution

Submit your manuscript at www.biomedcentral.com/submit