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Classic axon guidance molecules control correct nerve bridge tissue formation and precise axon regeneration

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Abstract

The peripheral nervous system has an astonishing ability to regenerate following a compression or crush injury; however, the potential for full repair following a transaction injury is much less. Currently, the major clinical challenge for peripheral nerve repair come from long gaps between the proximal and distal nerve stumps, which prevent regenerating axons reaching the distal nerve. Precise axon targeting during nervous system development is controlled by families of axon guidance molecules including Netrins, Slits, Ephrins and Semaphorins. Several recent studies have indicated key roles of Netrin1, Slit3 and EphrinB2 signalling in controlling the formation of new nerve bridge tissue and precise axon regeneration after peripheral nerve transection injury. Inside the nerve bridge, nerve fibroblasts express EphrinB2 while migrating Schwann cells express the receptor EphB2. EphrinB2/EphB2 signalling between nerve fibroblasts and migrating Schwann cells is required for Sox2 upregulation in Schwann cells and the formation of Schwann cell cords within the nerve bridge to allow directional axon growth to the distal nerve stump. Macrophages in the outermost layer of the nerve bridge express Slit3 while migrating Schwann cells and regenerating axons express the receptor Robo1; within Schwann cells, Robo1 expression is also Sox2-dependent. Slit3/Robo1 signalling is required to keep migrating Schwann cells and regenerating axons inside the nerve bridge. In addition to the Slit3/Robo1 signalling system, migrating Schwann cells also express Netrin1 and regenerating axons express the DCC receptor. It appears that migrating Schwann cells could also use Netrin1 as a guidance cue to direct regenerating axons across the peripheral nerve gap. Engineered neural tissues have been suggested as promising alternatives for the repair of large peripheral nerve gaps. Therefore, understanding the function of classic axon guidance molecules in nerve bridge formation and their roles in axon regeneration could be highly beneficial in developing engineered neural tissue for more effective peripheral nerve repair.

Key Words: axonal guidance; EphrinB2; nerve bridge; Netrin1; peripheral nerve; regeneration; Slit3; Sox2; transaction injury

Introduction

Injuries to peripheral nerves are common and often result in the complete transaction of the nerve (Pfister et al., 2011). Such an injury may create a gap between the proximal and distal nerve stumps, requiring either an autologous nerve graft or the use of an artificial nerve conduit for repair when the gap is greater than 5 mm (Moore et al., 2009). Studies using a sciatic nerve transection injury model in mice and rats have shown that new nerve bridge tissue can form between the proximal and the distal nerve ends and that regenerating axons are able to navigate across the nerve gap using the newly-formed bridge tissue as a substrate (Parrinello et al., 2010; Cattin et al., 2015; Dun and Parkinson, 2015; Dun et al., 2019). Engineered neural tissues have been suggested as promising alternatives for the repair of large peripheral nerve gaps to replace the current gold standard of using nerve autografts (Georgiou et al., 2013). Therefore, understanding the molecular mechanism of nerve bridge tissue formation is crucial not only to the understanding of basic principles governing cell biology and tissue regeneration, but also to the development of novel engineered neural tissue for effective nerve repair. Previous studies on roles for Netrin1/DCC signaling and EphrinB2/EphB2 signaling (Parrinello et al., 2010; Webber et al., 2011; Rosenberg et al., 2014), and more recently our study on roles for Slit3/Robo1 signalling within the nerve bridge, have shown that classic axon guidance molecules play key roles in regulating Schwann cell migration and precise axon regeneration in the peripheral nerve bridge (Dun et al., 2019). In this review, we will summarize the current understanding of how classic axon guidance molecules regulate cell migration, nerve bridge formation and precise axon regeneration in the peripheral nerve gap after transaction injury (Figure 1).

The current scientific literature was searched using PubMed with the keywords Netrin, Slit, Ephrin or Semaphorin together with the key words peripheral, nerve and regeneration. The search for ‘Netrin’ retrieved 17 published papers; the search for ‘Slit’ retrieved 21 published papers; the search for ‘Ephrin’ retrieved 14 published papers and the search for ‘Semaphorin’ retrieved 28 published papers. Papers were then screened and assessed for inclusion according to their significance for the field of signalling in the formation of nerve bridge tissue following injury.

Netrin1/DCC Signaling between Migrating Schwann Cells and Regenerating Axons Directs Regenerating Axons Across the Nerve Bridge

Netrins are a family of extracellular, laminin-related proteins that are widely expressed in a range of embryonic tissues and play key roles in axonal guidance during nervous system...
development (Dun and Parkinson, 2017). In mammals, four secreted Netrins, Netrin1, 3, 4 and 5, have been identified, and so far Netrin1 is the best characterized member of the family (Lai Wing Sun et al., 2011; Yamagishi et al., 2015). Netrin1 was the first axon guidance molecule to be discovered in 1994, representing a true milestone in the research field of axonal guidance (Kennedy et al., 1994; Serafini et al., 1994). It can bind to a variety of receptors such as DCC, Neogenin, Unc5A-D and CD146 to transduce receptor-specific biological functions but its attractive function for axon extension is largely transduced by the DCC receptor (Dominici et al., 2017; Dun and Parkinson, 2017). Netrin1 was shown to be up-regulated after peripheral nerve injury and the expression was primarily in Schwann cells (Madison et al., 2000; Petrausch et al., 2000; Webber et al., 2011; Jaminet et al., 2013; Rosenberg et al., 2014). Studies have shown that Netrin1 plays a vital role in adult peripheral nerve regeneration (Webber et al., 2011; Jaminet et al., 2013; Rosenberg et al., 2014; Ke et al., 2015). Netrin1 heterozygous mice showed a much slower functional recovery after peripheral nerve crush injury and full functional recovery was not observed even at 50 days following injury (Jaminet et al., 2013). Injection of Netrin1 overexpressing bone marrow mesenchymal stem cells into the crush site of rat sciatic nerve promoted more rapid axon regeneration and functional recovery (Ke et al., 2015). In the zebra fish mutant deleting DCC in motor neurons, regenerating motor axons in the nerve bridge strayed away from their original path onto ectopic trajectories, recapitulating the phenotype observed in zebra fish mutant lacking migrating Schwann cells in the nerve bridge (Rosenberg et al., 2014). Binding of Netrin1 to the DCC receptor typically transduces Netrin1 attractive signaling (Dominici et al., 2017; Dun and Parkinson, 2017). Thus, these findings indicated that Schwann cell could utilise Netrin1 as a guidance cue to direct axon regeneration in peripheral nerve bridge via the DCC receptor expressed on regenerating axons.

**EphrinB2/EphB2 Signaling between Fibroblast and Migrating Schwann Cells Regulates Schwann Cell Cord Formation**

Eph receptors (erythropoietin-producing human hepatocellular receptors) belong to the largest family of tyrosine kinase receptors and they are activated upon the binding of their ligands, the Eph receptor-interacting proteins, Ephrins (Lisabeth et al., 2013). Ephrin/Eph signaling not only has a critical function in embryonic nervous system development by regulating axonal guidance and cell migration but also plays an important role in tissue repair following injury (Coulthard et al., 2012). Previously, Parrinello et al. (2010) showed that EphrinB2/EphB2 signaling between fibroblasts and Schwann cells is critical for the formation of Schwann cell cords within the nerve bridge during regeneration. In the nerve bridge, migrating Schwann cells are surrounded and contacted by nerve fibroblasts but the two cell types do not appear to mix with each other. In co-culture experiments, Schwann cells and nerve fibroblasts sorted into mutually exclusive cell clusters. Examining nerve bridge tissue as well as cultured nerve fibroblasts and Schwann cells, Parrinello et al. (2010) demonstrated that Schwann cells express the EphB2 receptor while nerve fibroblasts express the EphrinB2 ligand, suggesting that EphrinB2/EphB2 signaling may be required for the coordination between nerve fibroblasts and Schwann cells in the nerve bridge. By manipulating the levels of EphrinB2 and EphB2 in cell co-cultures, Parrinello et al. (2010) demonstrated that EphrinB2/EphB2 signaling between nerve fibroblasts and Schwann cells is necessary and sufficient for Schwann cell sorting and cluster formation. This study also examined axon regeneration in the nerve bridge of EphB2 knockout mice or in wild-type rats in which an inhibitory EphB2-Fc fusion protein was delivered to the nerve bridge via mini osmotic pumps. In both cases, axonal regrowth appeared less organized in the nerve bridge compared to control experiments. As regenerating axons are guided by the Schwann cell cords in the nerve bridge, these findings demonstrated that EphrinB2/EphB2 signaling between nerve fibroblasts and Schwann cells in the nerve bridge is important for correct nerve bridge tissue formation.

Although Ephrin/Eph signaling is thought to primarily induce rapid cell responses by controlling actin dynamics, Parrinello et al. (2010) found that increased EphrinB2/EphB2 signaling increases the expression of the transcription factor Sox2 in Schwann cells. Sox2 expression in Schwann cells promotes tissue repair under several different circumstances including peripheral nerve bridge formation, skin wound healing and digit tip regeneration (Johnston et al., 2016; Carr and Johnston, 2017; Parfejevs et al., 2018). Parrinello et al. (2010) showed that EphrinB2/EphB2 signaling via Sox2 promotes the translocation of the cell surface adhesion molecule N-cadherin into the Schwann cell membrane. This increased level of N-cadherin promoted adhesion between Schwann cells. By knockdown of EphB2, Sox2 and N-cadherin levels in cultured Schwann cells, Parrinello et al. (2010) showed that N-cadherin is necessary and sufficient for cell sorting downstream of EphB2 and Sox2 in vitro. Sox2 overexpression rescued the Schwann cell sorting deficiency of cultured Schwann cells from EphB2 knockout mice, indicating that Sox2 acts downstream of EphB2 to regulate Schwann cell sorting. Thus, these findings uncovered a new signaling pathway downstream of EphB2 activation via Sox2 and N-cadherin that mediated the Schwann cell sorting process in the nerve bridge, ultimately leading to the formation of Schwann cell cords in the nerve bridge.

**Macrophage-Derived Slit3 Controls Correct Nerve Bridge Tissue Formation**

In another recent paper, we examined the pattern of axon regeneration and the trajectory of Schwann cell migration in the nerve bridge of mice with a Schwann cell-specific knockout of Sox2 (Dun et al., 2019). We showed a dramatic defect of both axon pathfinding and Schwann cell migration in the sciatic nerve bridge of Sox2 knockout mice following nerve injury. At both 10 and 14 days following sciatic nerve transection, large numbers of axons had left the nerve bridge and a completely abnormal nerve bridge formation was still clearly observed even at three months post-injury. To test whether the axon regeneration defects in Sox2 knockout mice were caused by ectopic Schwann cell migration, we used green fluorescent protein-labelled Schwann cells by crossing Sox2 knockout mice with a proteolipid protein promoter driving green fluorescent protein line (Mallon et al., 2002). Abnormal Schwann cell migration within the nerve bridge is the first step in Sox2 knockout mice, indicating that Sox2 acts downstream of EphB2 to regulate Schwann cell sorting. Thus, these findings uncovered a new signaling pathway downstream of EphB2 activation via Sox2 and N-cadherin that mediated the Schwann cell sorting process in the nerve bridge, ultimately leading to the formation of Schwann cell cords in the nerve bridge.

bridge of Sox2 knockout animals was observed at 6 days following sciatic nerve transection with regenerating axons following the ectopic migrating Schwann cells. Ectopic migrating Schwann cells in Sox2 knockout nerves were still observed at 14 days post-injury and did not form correct Schwann cell cords connecting the proximal and the distal nerve stumps. To examine potential Sox2 targets in Schwann cells, we performed in vitro microarray analysis by overexpressing Sox2 in cultured primary rat Schwann cells and identified that Sox2 also regulates the expression of the axon guidance receptor Robo1. The regulation of Robo1 expression by Sox2 in Schwann cells was further confirmed in both Schwann cell-specific Sox2 knockout and Sox2 overexpressing mice after sciatic nerve transection injury (Dun et al., 2019).

The ligands for Robo1 are the secreted Slit glycoproteins. So far, three Slits (Slit1/2/3) have been identified in vertebrates with a spatio-temporal expression pattern in the central and peripheral nervous system during development and all of them bind to the Robo1 and Robo2 receptors with high affinity (Blockus and Chedotal, 2016). Interaction of Slit1–3 ligands with their Robo1–2 receptors form one of the most crucial ligand-receptor pairs among the classic axon guidance molecule family proteins and serves as a repulsive signaling pathway to control precise axon pathfinding and neuronal migration during nervous system development (Blockus and Chedotal, 2016). This is a function that is conserved in flies, worms, and vertebrates and which has been validated in numerous genetic studies (Blockus and Chedotal, 2016). After identifying that Sox2 regulates Robo1 expression in Schwann cells, we moved on to investigate the expression of Slit1–3 and Robo1–2 in the nerve bridge and found that Slit3 and Robo1 are highly expressed in the mouse sciatric nerve bridge during regeneration (Dun et al., 2019). Macrophages in the outermost layer of the nerve bridge express high levels of Slit3 while migrating Schwann cells inside the nerve bridge express high levels of its receptor Robo1. The expression of Slit3 by macrophages in the outermost layer of the nerve bridge and the expression of Robo1 in migrating Schwann cells inside the nerve bridge, led us to speculate that macrophage derived Slit3 could regulate the trajectory of Robo1 expressing Schwann cell migration within the nerve bridge.

To understand a primary role of Slit3/Robo1 signaling in Schwann cell migration, mice with reduced expression of both Slit3 and Robo1 (Slit3+/−/Robo1+/−) were generated and the effects upon Schwann cell migration and axonal pathfinding were tested. In control animals, at 14 days post-transection, Schwann cells migrated and stayed within the bridge and formed cell cords connecting the proximal and distal nerve ends. However, in Slit3+/−/Robo1−/− mice, a large population of Schwann cells left the nerve bridge from both the proximal and the distal nerve ends, confirming that Slit3 is required for controlling the trajectory of Schwann cell migration within the nerve bridge. We also co-cultured rat primary Schwann cells with control Slit3+/− or Slit3−/− bone marrow macrophages and demonstrated an apparent Slit3/Robo1 repulsive signaling between Schwann cells and macrophages. A further examination of nerve bridge formation confirmed that the nerve bridge tissue was not correctly formed in Slit3+/−/Robo1−/− mice. Further studies have been carried out to examine the pattern of axon regeneration in the nerve bridge of Slit1−3 and Robo1−2 gene mutant mice at 14 days following sciatic nerve transection injury. In line with the observation of ectopic Schwann cell migration in Slit3 and Robo1 gene mutant mice, a remarkable axon regeneration defect was observed in the nerve bridge of Slit3+/β2/Robo1−/− mice but not in Slit1−/−, Slit2−/−, or Robo2−/− mice. Thus, our studies have demonstrated that the Slit3/Robo1 repulsive signaling between macrophages...
and Schwann cells in the nerve bridge plays an important role in regulating the trajectory of Schwann cell migration. Angiogenesis is a key event for correct nerve bridge formation and all classic axon guidance molecules have important functions in angiogenesis (Adams and Eichmann, 2010). However, we found that Slit3/Robo1 signaling does not appear to be required for angiogenesis in the nerve bridge following nerve transection (Dun et al., 2019). It will be interesting to study if other classic axonal guidance signaling pathways are involved in regulating angiogenesis in the nerve bridge.

Future Direction

Successful peripheral nerve bridge formation requires the coordination of multiple cell types and the activation of multiple signaling pathways in the nerve bridge. The nerve bridge consists of newly formed tissue connecting the proximal and distal nerve stumps and comprises macrophages, fibroblasts, endothelial cells and Schwann cells (Cattin et al., 2015). Previously, Parrinello et al. (2010) showed that the coordination between nerve fibroblasts and Schwann cells is controlled by EphrinB2/EphB2 signaling, and recently we showed that the coordination between macrophages and Schwann cells is controlled by the Slit3/Robo1 signaling. During development, classic axon guidance molecules Netrin, Slit, Ephrin and Semaphorin often cooperate with each other to regulate cell migration and axon pathfinding (Bashaw and Klein, 2010). Cell coordination in the nerve bridge between perineurial cells and Schwann cells (Lewis and Kucenas, 2014), between macrophages and endothelial cells (Cattin et al., 2015), and between macrophages and fibroblasts (Dun et al., 2019) have all now been reported. It will be interesting to study how other axonal guidance signaling pathways are involved in controlling cell coordination in the nerve bridge, and how the synergistic function among classic axon guidance molecules controls correct nerve bridge tissue formation and precise axon targeting. We believe understanding these key cellular and molecular mechanisms could help to develop better engineered neural tissue for more effective peripheral nerve repair.

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