Recurrent laryngeal nerve (RLN) injury and voice alteration after thyroid surgery

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Abstract: Loss of function in the recurrent laryngeal nerve (RLN) during thyroid/parathyroid surgery, despite a macroscopically intact nerve, is a challenge which highlights the sensitivity and complexity of laryngeal innervation. Furthermore, the uncertain prognosis stresses a lack of capability to diagnose the reason behind the impaired function. There is a great deal of literature considering risk factors, surgical technique and mechanisms outside the nerve affecting the incidence of RLN paresis during surgery. To be able to prognosticate recovery in cases of laryngeal dysfunction and voice changes after thyroid surgery, the surgeon would first need to define the presence, location, and type of laryngeal nerve injury. There is little data describing the events within the nerve and the neurobiological reasons for the impaired function related to potential recovery and prognosis. In addition, very little data has been presented in order to clarify any differences between the transient and permanent injury of the RLN. This review aims, from an anatomical and neurobiological perspective, to provide an update on the current understandings of surgically-induced injury to the laryngeal nerves.

Keywords: Regeneration; thyroid surgery; nerve injury; vocal fold paresis; laryngeal EMG

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Thyroid and parathyroid surgery is associated with a risk of traumatic injury to the superior and inferior RLN. Injury to the RLN results in acute paralysis of the vocal fold on the affected side, which leads to dysphonia, dysphagia, and aspiration problems. The clinical signs may vary, however, depending on the position of the paralyzed vocal fold relative to the midline and the degree of glottic insufficiency. A significant number of patients may even present as asymptomatic, and vocal fold mobility needs to be examined before and after surgery to detect an iatrogenic injury to the RLN (1-4). The reported risk for RLN injury after thyroid/parathyroid surgery varies from center to center. In the literature, injury rates of up to 38% can be found (5). It is difficult to compare these numbers, since the rate of detected RLN injury is dependent on how close to surgery the laryngoscopic examination is performed. In a large, retrospective study of patients that underwent total thyroidectomy due to malignant disease, the risk of postoperative vocal fold paresis was found to be 9.5% (6), of which 22% became permanent with resulting demand for secondary surgical intervention of the paralyzed vocal fold.

Subjective voice deficits are common after thyroid surgery (7). Subjective and objective voice alterations (8,9) are common after total thyroidectomy and most of them are independent of injury to the RLN or the superior laryngeal nerve (SLN) (10-12). These voice changes are believed to be caused by temporary disturbances in the laryngeal mechanical framework or extralaryngeal scarring, and they usually resolve to subclinical levels within weeks or months after the operation (9,13,14). Dividing the sternothyroid muscle has been shown not to affect voice outcome in a significant way (15). Even though the RLN is the most important provider of laryngeal motor innervation, injury
to the external branch of the SLN is also believed to cause significant voice changes, such as reduction in the fundamental frequency range, reduction in the highest obtainable fundamental frequency and vibratory phase asymmetry in the vocal folds (16-19). The external branch of the SLN has therefore gained increased attention among thyroid surgeons, and it is recommended that it should be preserved as well as the RLN during thyroid surgery (20).

In most cases of postoperative vocal fold paresis, the RLN is macroscopically intact and the injury is located within the peripheral nerve. To be able to prognosticate recovery, the surgeon would need to define the presence, location, and type of the nerve injury. However, very little data has been presented aimed at clarifying the differences between the transient and permanent injury of the RLN. This review aims to provide, from an anatomical and neurobiological perspective, an update on the current understanding of surgically-induced injury to the laryngeal nerves.

The laryngeal nerves

The larynx is innervated by two branches of the vagus nerve, the RLN, and the SLN. Both nerves carry motor, sensory, and autonomic (parasympathetic) fibers to the larynx (21). The lower motor neurons of the special efferent system controlling the intrinsic laryngeal muscles are located in the nucleus ambiguus in the lower brainstem, in a fairly well-studied somatotopical arrangement (22-30). The sensory neurons are located in the nodose ganglion and the parasympathetic cell bodies are located in the dorsal motor nucleus of the vagus in the brainstem. The mechanically complex laryngeal functions (airway protection reflexes, phonation, swallowing) require a rich and detailed neural control, projected through the RLN and SLN.

The RLN can be regarded as the most important motor nerve supply to the larynx, as it innervates 4 out of 5 intrinsic laryngeal muscles. It also has projections to the esophagus and trachea (23). The RLN divides into an anterior and posterior branch. The branching point is located either inside the larynx, or, as in roughly one third of cases, before the nerve entering point (31). The posterior branch projects superiorly to form the anastomosis of Galen with the internal branch of the SLN—this branch is probably sensory in nature (32). The anterior branch carries motor fibers (33,34) to the posterior thyroarytenoid muscle, lateral cricoarytenoid muscle and, finally, to the thyroarytenoid muscle (35-37). Within the larynx, the RLN and SLN break up into a plexus-like branching system (38), with several connections between the RLN and SLN (37). The exact functions of these small nerve branches are not fully known, but it has been demonstrated in animal experimental models that the intrinsic laryngeal muscles receive dual innervation from both the RLN and SLN (39,40).

The SLN originates from the inferior vagal ganglion at the C2 level in the neck (41). It divides into a larger, internal branch which enters the larynx through the thyrohyoid membrane (carrying sensory fibers down to the level of the glottis) and a smaller, external, branch which passes deep to the superior thyroid artery to innervate the cricothyroid muscle responsible for vocal fold lengthening and tension, important for high voice pitch (42). The external branch of the SLN continues through the cricothyroid muscle to reach the anterior glottis and the thyroarytenoid muscle. This branch, called “the human communicating nerve” (43), or “the cricothyroid connection branch” (44) thus represents an additional motor supply to the intrinsic laryngeal muscles other than the RLN, which may be important following RLN injury and reinnervation. This anatomy enables intraoperative monitoring of the external branch of the SLN through routine surface electrodes in the intubation tube (45), although the exact laryngological function of this nerve branch is not known (17).

Neurapraxia versus axonotmesis

From a clinical perspective, it is important to make the distinction between nerve conduction block, “neurapraxia”, and the more severe “axonotmesis”, which means presence of axonal injury (Figure 1). These classifications were first made by Seddon in 1942 (46) and later modified by Sunderland in 1951 (47). Surgically-induced nerve injuries seldom include complete transection of the nerve, but rather intraneural damage inside a macroscopically intact nerve due to pressure, crush or heating from adjacent use of cautery. Neurapraxia is the mildest form of injury, affecting the surrounding Schwann cells, but respecting the integrity of the axon (Figure 1A). The result is a conduction block lasting typically about 6-8 weeks followed by a complete return of function, when the Schwann cells have been repaired (48). This seems to be the case also for RLN injury (49). Following axonotmesis, there is a varying degree of axonal injury (Figure 1B), which could lead to neuronal death or dysfunctional reinnervation of the target cells. Axonotmesis, therefore, is associated with a poorer and more unpredictable
outcome for functional restitution.

**Neurobiology behind impaired function**

The motor neuron terminates at the neuromuscular junction, the motor end plate. The neuron is the secondary neuron and is part of the peripheral nervous system (PNS), as opposed to the primary motor neuron [central nervous system (CNS)], which runs from the cortex to terminate on the secondary neurons. The myelin around the axons in the CNS comes from oligodendrocytes. The CNS myelin contains several factors which are inhibitory to axonal growth and regeneration, which is one of the major problems after CNS injury, such as stroke or spinal cord injury (50). In the PNS, on the other hand, the myelin around the axons is derived from the Schwann cell. This milieu is attractive for axonal growth, which is why the peripheral nerve injury is usually associated with regeneration after axonal disruption (51-55). Following a peripheral nerve injury, which involves peripheral axon disruption, the distal part (which is disconnected from the Schwann cell) will be neurophysiologically active until it degenerates (56) [Wallerian degeneration (57)] which, under normal conditions, will take approximately one week. Thus, a complete injury to the RLN which separates the nerve into two different parts, will give a negative signal using intraoperative nerve monitoring (IONM) and the distal part a positive signal in the thyroarytenoid muscle for several days. Re-exploring the distal end of the RLN at our institution for nerve re-construction confirms the positive signal for up to five days after complete injury after thyroid surgery (unpublished observation). After axonal injury within the macroscopically intact RLN, the distal axon also degenerates and, to achieve any functional recovery, the axon has to regenerate. In the literature, there has been considerable speculation concerning the reasons for the poor (or absent) functional recovery seen after injury to the RLN despite the fact that the nerve looks macroscopically intact during surgery. One factor associated with the insufficient recovery is a potential misguidance of RLN axons during regeneration, leading to non-functional reinnervation of laryngeal muscles. There are, however, studies that show that the degree of accurate innervation is very high after crush injury to the peripheral nerve (90%) (58) since the axon is guided by intact mechanical factors of the intact endoneural tubes (59). This may reduce the impact of the misguidance as a negative factor in the injured intact RLN.

The axotomy induces a retrograde injury signal to the neuron in the brainstem which is attacked by microglia and also surrounded by a profound astroglial reaction (60-62). The neuron downregulates its production of transmitter substances and turns the gene transcription to regeneration and re-innervation. The neuron is exposed to stress and is dependent on a continuous inflow of growth factors from the periphery (54). Motor neurons are more likely to die in response to peripheral axotomy the closer the axotomy is to the neuronal soma in the brainstem or spinal cord. The

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**Figure 1** Schematic drawing of the RLN with intralaryngeal branches to IA, LCA and TA. The axon is surrounded by Schwann cells responsible for electrical propagation. Neurapraxia (A) with intact axonal integrity, facing spontaneous recovery. Axonotmesis (B) with disruption of axon and ongoing regeneration. RLN, recurrent laryngeal nerve; IA, interarytenoid muscle; LCA, lateral cricoarytenoid muscle; TA, thyroarytenoid muscle; LB, ligament of Berry; GA, anastomosis of Galen.
more of the peripheral nerve which is in contact with the neuron soma, the more trophic support of growth factors is delivered to the neuron. The addition of growth factors radically improves the prognosis for the axotomized neuron (63-66). In addition, there are many experiments to support that distal peripheral nerve injury is associated with no or limited nerve cell death, including injury to the RLN (67).

The distal axotomy in the intact RLN also causes a synaptic displacement from the secondary motor neuron in the nucleus ambiguous, which then loses contact with higher cortical centers (68). These synapses from cortical neurons re-appear on the secondary motor neurons as the neurons manage to regenerate and re-establish contact with the target organ (muscle). The proceeding adaptation to the new neural circuits is referred to as plasticity of the nervous system. Thus, the macroscopically intact but injured RLN will recover spontaneously if there is only a conduction block caused by an impairment of electrical propagation due to Schwann cell affection. If there is a component of axonal injury within the nerve, the axon will not only have to re-innervate the laryngeal muscles, but the neuron in the brainstem will need a re-connection with cortical neurons by re-establishment of their synapses onto the secondary neuron in the brainstem.

**Intraoperative nerve monitoring (IONM)**

IONM of the RLN is performed by stimulating the peripheral nerve directly with an electrical current, with subsequent recording of muscle depolarization of the thyroarytenoid muscle. When a peripheral nerve is directly stimulated at supramaximal intensity, the result is depolarization of all axons and activation of all motor units projecting through the nerve, which leads to acetylcholine-mediated depolarization of muscle fibers. The intramuscular shift in electrical potential (voltage) can be recorded as a compound muscle action potential (CMAP), representing the sum of all motor unit activity. During IONM, the presence and the amplitude of the CMAP is utilized as an indirect measurement of motor nerve function during the surgery. Originally, RLN monitoring was made through needle electrodes inserted into the intrinsic laryngeal muscles. In modern clinical routine, thyroarytenoid depolarization is recorded through surface electrodes on the ventilation tube (69).

Manipulation of the surgical field may affect the RLN by traction, heating, entrapment or squeezing (crush injury), which leads to absence or reduced amplitude of the recorded CMAP following vagal stimulation. The neurobiological explanation for reduced CMAP amplitude is simply that a lower number of axons are transmitting the electrical signal, which means less depolarization of the monitored muscle. Provided that the vagal stimulation is performed in the same way [consistent with continuous monitoring (70)], the site of injury could be anywhere along the nerve, distal to the site of stimulation. When the signal is lost during surgery, it is not possible to use IONM to diagnose the type of nerve injury (axonotmesis or neurapraxia), it only tells us that there is a discontinuity of the electrical propagation within the nerve.

In order to categorize and group injuries to the RLN using IONM, one reported way is to define the injury to the RLN and loss of signal (LOS) from the vagal nerve as segmental (type 1) or global (type 2) (69,71). Looking at the basic neurobiology of the nerve, a LOS could originate from an injury anywhere from the stimulus (vagal nerve) to the neuromuscular endplate, including the muscle. The distal part of the nerve is excitable for several days after injury even after nerve transaction injury (48), which makes it possible for the surgeon to pin-point the exact location of the nerve injury, by using the stimulation probe along the course of the peripheral nerve. A neurobiological explanation for the “global” (type 2) RLN injury could be that the location of the nerve injury is located distal to the nerve entry point under the inferior constrictor muscle, not affecting the whole neuron (i.e., a milder form of injury). From this perspective, type 1 and 2 injuries describe if the RLN conduction block is proximal or distal to the RLN/cricothyroid border (Figure 2).

**Laryngeal EMG**

Postoperative electrodiagnostic methods can be used to determine the presence and type of nerve injury, as well as to characterize the ongoing or completed reinnervation processes. Laryngeal electromyography (LEMG) was first introduced more than sixty years ago, and has evolved (72) into a valuable tool for laryngologists in diagnosing neurolaryngological disorders. It has been pointed out that LEMG is primarily a qualitative method (73) (presence of denervation potentials, degree of motor unit recruitment), which makes it a subjective test depending on the examiner and the technical settings. But LEMG has nevertheless been shown to have high positive predictive value in predicting the long-term outcome of patients with a poor prognosis (74-77) and it is used widely to predict...
recovery regardless of the etiology behind the vocal fold paresis. Patients with pathological electromyographic findings at least two months after the paresis are most likely to need laryngeal framework surgery (76). In the case of postoperative vocal fold paresis after thyroid/parathyroid surgery, the prognostic information obtained from LEMG can be helpful to identify those cases where future interventions are necessary, which could mean surgical or pharmacological reinnervation therapies, or vocal fold medialization procedures. For patients with only a conduction block (neurapraxia) of the RLN, vocal fold movement is most likely to return. When using LEMG to obtain this information after thyroid surgery, it is important to take into consideration the timing of the examination. Denervation activity (indicating axonotmesis and poor prognosis) typically appears at three weeks after the RLN injury (48), and lasts until reinnervation is complete. Reinnervation of the intrinsic laryngeal muscles can be expected to take place rather promptly, given the high regenerative capacity of the RLN (78), together with collateral reinnervation by adjacent, intact nerve fibers (40). The optimal time window for postoperative LEMG seems to be 2-4 weeks after the nerve injury (49). Interpretation and analysis of electrophysiological data requires the expertise of a trained neurologist or clinical neurophysiologist, while insertion of the needle electrodes into the appropriate intrinsic laryngeal muscles is best performed by an ENT specialist. LEMG thus requires the cooperation and coordination of different clinical resources. A consensus paper for LEMG guidelines in the areas of indications, technical considerations, implementations and data interpretation was published by Volk et al. (79) in 2012.

**Regeneration promotion**

A mixed injury of demyelination (neurapraxia) and axonotmesis within the macroscopically intact RLN has a worse prognosis than demyelination alone, because of the need for regeneration and reinnervation of the target. Reinnervation of the intrinsic laryngeal muscles following axonotmesis is considered problematic (80), due to misguided, unordered regeneration and perhaps also collateral reinnervation originating from adjacent, intact nerve fibers (40). Pathological reinnervation leads to a change in the somatotopic map, not in line with normal vocal fold function. Theoretically, it would be beneficial for the functional restitution to pace up and enhance regeneration/reinnervation by the RLN.

*In vitro*, it has been shown that the pace of the regenerating axon is regulated at the tip (growth cone), the motion of which is highly dependent on a delicate regulation of calcium ions (81,82). It was demonstrated that altering the intracellular concentration of calcium ions had a strict correlation to the ability of the growth cone to sprout (82). The regulation of intracellular calcium is also closely linked to the actions of the voltage-gated calcium channels present in the cellular membrane (82). *In vivo*, it has been confirmed that the transient quick calcium currents across the membrane of the growth cone occur with a certain frequency. If the calcium transient calcium currents are to some extent inhibited the pace of axonal elongation increases, and vice versa (83-85). In fact, blocking of the rapid calcium flow current across the membrane would increase the total time for axonal elongation, a principle further evaluated in experimental models. Nimodipine, a voltage-gated calcium flow antagonist to the L-type channels has been evaluated in rodent models, and is a pharmacologically good choice because it penetrates the blood brain barrier better than most other calcium flow antagonists (86). After systemic administration of

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**Figure 2** The RLN and the NEP under the IC. NEP may serve as an anatomical landmark in the classification of loss of vagal signal during thyroid surgery, e.g., lesion proximal to NEP (when there is a defined injury segment) or lesion distal to NEP (silent nerve to the NEP). RLN, recurrent laryngeal nerve; NEP, nerve entry point; IC, inferior constrictor muscle; TG, thyroid gland; SLNi, superior laryngeal nerve internal branch; GA, anastomosis of Galen; LB, ligament of Berry.
nimodipine, an improved regeneration and functional recovery has experimentally been achieved after injury to the sciatic (87), facial (88-90), hypoglossal (91) and RLNs (92). In the patient, nimodipine has been evaluated after recurrent laryngeal (49,93-95) and facial nerve injury (96-100), with promising functional outcomes. Taken together, there is substantial evidence that the administration of nimodipine after axonal injury to a peripheral nerve probably improves the functional outcome.

Even though there is emerging data that treatment with nimodipine may also be translated to the patient in some situations, the level of evidence for a using nimodipine for intraoperative RLN injury is still modest. Only a fraction of the patients with postoperative RLN paresis would benefit from a regeneration-promoting treatment (i.e., cases with axonotmesis). It is important, therefore, to search for further knowledge concerning diagnosis and prognosis after RLN injury after thyroid surgery.

Conclusions

Laryngeal dysfunction and voice problems are common after thyroid surgery, but only a fraction of these cases turn out to be chronic. Chronic laryngeal dysfunction is most commonly caused by axonal injury to the RLN or SLN. The clinical progress of symptoms and the eventual functional recovery of the target organ follow the general principles of peripheral nerve injury, even though the larynx can be regarded a special case in being functionally and neuroanatomically complex, with high demands for accurate neural supply. Today, it is possible for the clinician to utilize the information obtained from electrodiagnostic methods (IONM and postoperative LEMG), to characterize the nerve injury and predict the temporal course and functional result of the healing process. It is important to do so in order to be prepared for additional interventions, such as voice therapy, medialization surgery or regeneration/reinnervation therapies.

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