Review Article

Experimental spinal cord injury and behavioral tests in laboratory rats

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Abstract

Traumatic spinal cord injury (SCI) results in some serious neurophysiological consequences that alter healthy body functions and devastate the quality of living of individuals. To find a cure for SCI, researchers around the world are working on different neurorepair and neurorehabilitation modalities. To test a new treatment for SCI as well as to understand the mechanism of recovery, animal models are being widely used. Among them, SCI rat models are arguably the most prominent. Furthermore, it is important to select a suitable behavioral test to evaluate both the motor and sensory recovery following any therapeutic intervention. In this paper, we review the rat models of spinal injury and commonly used behavioral tests to serve as a useful guideline for neuroscientists in the field of SCI research.

Keywords: Neurology, Physiology, Rehabilitation, Surgery, Anatomy, Neuroscience

1. Introduction

Spinal cord injury (SCI) is responsible for damaging the sensory, motor and autonomic functions of an individual [1]. To address this problem, over the past few
decades animals have been used for developing and testing therapies for SCI [2]. Both large and small animal models are commonly used to study the injury mechanism and to assess the therapeutic outcomes after a controlled SCI in laboratory settings [3]. Large animals, especially dogs and cats were used in the early 1950s—1970s [4]. Afterward, rodent models became popular and were then standardized as a suitable animal model for SCI experiments. In rodent models, rats and mice are mostly used to evaluate the functional and morphological changes after SCI [5, 6, 7]. However, in the mouse model, following a SCI, evidence suggests that a modest degree of regeneration occurs in the injured area and no typical cyst-like structure is formed, while such type of axonal regeneration is not found in rat and non-human primates [1]. In contrast, after an injury in rats, a large fluid-filled cystic cavity is formed which is similar to the human pathology after SCI and thus useful for preclinical studies [8]. Rats are quickly adaptive to the experimental environments and to the given tasks for sensorimotor evaluations. Furthermore, high availability, low infection rate after surgery, and ease of maintenance make the rat a prime candidate for translational research. Based on the above, laboratory rats are now widely used as experimental SCI models.

In this translational research, several rat injury models were developed to examine the neurophysiological responses after a SCI. Numerous approaches are being applied to regenerate the damaged axon across the lesioned area, and confirmed immunohistochemically and by protein analysis in rats [9, 10]. Furthermore, apart from histological analysis, behavioral responses are also studied as a primary measure to assess the recovery of spinal pathways. Other reviews [11, 12, 13, 14] focused on the different animal models of SCI particularly cervical injury and some behavior tests. However, to our knowledge, no study has comprehensively described about all the injury models and their impact on the behavioral changes. In this review, we focused on different SCI rat models and their recovery assessments using different behavioral tests. We also propose some novel behavioral tests to enrich this research. Our review offers a guideline for the researchers in this field to successfully conduct SCI research using rats.

2. Main text

2.1. Rat SCI models

Experimental injury models are designed to explore the physiological changes after SCI to evaluate the treatments to repair the broken spinal cord. Laboratory rats are commonly used to evaluate neural repair after an experimental SCI. Although some SCI experiments have been conducted in non-human primates, more demanding protocols, critical manpower, health safety and a large amount of investment in the experimental setup and management make the primate model much more
difficult and less used than rat injury models [15]. In rat SCI experiments, three different injury models are widely utilized: compression, contusion and transection models. Among these, contusion and compression are the most common types of injury found in humans. Hence, to evaluate the neuronal changes and behavioral outcomes, these models are essential [7].

2.1.1. Compression model

In humans, flexion and axial compression (burst fracture) are responsible for displacing bone fragments into the vertebral canal and consequently SCI occurs. However, the timing of compression can affect the outcome of pathology [16]. To address this problem, in rats the same model of injury is carried out to create persistent spinal canal occlusion that is similar to human acute SCI. This injury is varied at different levels of the spinal cord. For example, at lumbar level, compression injury impairs the ejaculation reflex in male rats [17]. There are several ways to create a compression model for rats such as calibrated forcep compression, clip compression and balloon compression [16]. However, the clip compression model at the thoracic level is much more correlated with the functional and histological outcome [18].

The acute clip compression model is the first non-transection model that was developed in 1978 [19] and is widely used for experimental injury in rats. It is an inexpensive method where a special clip is used to compress the spinal cord [20]. Under inhalation anesthesia, a laminectomy is performed at a desired level of the spine by retracting muscles and removing the spinous process and body of vertebrae. A modified aneurysm clip (Fig. 1) is then inserted extradurally and maintained for 60
sec and then the clip is removed to produce an acute injury [21]. Muscles and connective tissues are then sutured, and the skin is closed.

Due to the similarity to human traumatic SCI, the compression injury model is also suitable for translational research. In a chronic condition of a compression SCI model, glial scars are formed which are quite similar to those in SCI patients [22]. To examine the therapeutic effect and neuroprotective study following a traumatic SCI, the compression model of injury is a better choice for the experiment. Furthermore, this model is useful to study the minimal loss of neurons after a SCI. In addition, the compression injury model is also useful to study the secondary injury mechanism events and cell transplantation therapies [23]. An important limitation of the compression method, however, is that after the injury some neural tracts may not be properly damaged which may lead to undesired outcomes in regeneration studies. Moreover, after 4 and 6 weeks of post-injury, tissue sparing may increase, suggesting that the late subacute and early intermediate stages of the secondary injury may be associated with limited neuroanatomical recovery [24].

### 2.1.2. Contusion model

In humans, compression injury is caused by mainly falls from a height or by a physical impactor that crushes the spinal vertebral column which further leads the bony parts to compress the spinal cord [1]. In rats, to get the same type of injury, the contusion model is probably the most suitable method [25]. Among the several anatomical locations, in thoracic position contusion injury is commonly used to evaluate the forward and backward locomotion as well as recovery of the spinal tract [26]. Usually, the injury is induced by an impactor which is used to drop a 10-gm rod onto the spinal cord [27]. Several impactors are available worldwide to serve this purpose. In Multicenter Animal Spinal Injury Study (MASCIS), the New York university impactor, Infinite Horizon (IH) impactor and Ohio State University Electromagnetic Spinal Cord Injury Device (ESCID) are most commonly used. In MASCIS, a 10-gm computer-controlled load is used to induce a SCI in rats [28]. After laminectomy, at a desired spinal level, muscles and spinous process are removed according to the diameter (usually 2.5 mm) of the impactor (Fig. 1). Height of the impactor, drop weight and time are previously set up on a computer before initiation of the injury. After the injury, subdural hemorrhage is commonly found which is later washed with saline [27]. Finally, muscles and skin are sutured. Another reliable commercially available impactor is Infinite Horizon (IH) which is also used to produce SCI rat models. System software is implemented to apply impactor force to the spinal cord. This injury renders an ideal model of SCI by reducing the variability among other existing devices [29].

Most SCIs reported in humans are similar to contusion and compression [8, 30]. Although the contusion SCI animal model is useful to study the secondary injury
mechanism and neuropathology of human injury [7], some limitations still persist [31]. In neuroregeneration research, after a thoracic contusion injury, it is difficult to study the fiber’s regeneration through the contusion site [32]. Furthermore, the contusion model cannot provide information regarding axonal tract regeneration which is essential to evaluate limb function [33].

2.1.3. Transection model

The transection model of injury is the most widely utilized method that can give the information of desired neural tracts at different spinal level after injury. To investigate the neuronal and functional improvements after different transplantation therapies, transection models of SCI are commonly used [34]. This injury model is quite stable and assessment can be performed within four weeks of the injury [35]. The spinal cord transection model is also useful to evaluate axonal regeneration and behavioral responses after SCI [36]. Transection injury cannot produce any condition similar to contusion injury. However, this injury is useful to investigate the damaged tract recovery after experimental therapies. Two main transection models (complete and incomplete) are used in rats. In the incomplete transection model (often called hemisection model), different portions of spinal cord are cut such as lateral hemisection, dorsal hemisection and dorsal funiculus crush to carry out different experiments. However, incomplete surgical transections are not considered relevant to human SCI [13]. A complete injury requires a complete transection of the spinal cord including both ascending and descending tracts. To carry out a complete transection injury after laminectomy, microdissection scissors are inserted to cut the spinal cord at a desired level and depth. After an injury gelfoam is inserted into the transection site to minimize the bleeding and to confirm the separation of the spinal cord. After some complete transections ventral parts of the axons may be spared, which is responsible for some residual motor function of the hindlimb [37]. For an incomplete transection injury iridectomy scissors are used to cut both the dorsal and ventral columns of the spinal cord from lateral to the midline by closing the tip of the scissors [38]. Muscles and skin incision are then sutured, and the rat is examined for therapeutic approaches and behavioral tests after recovery. Complete and incomplete transection injuries are not very similar to the clinical SCI of humans; they are rather utilized for neuroscience and neurology research to study the neuronal circuits and pathways [13].

Transection injury can give information on both ascending and descending tract functions which is useful in neuroscience research [7]. Compared to other injury models there is no need for a special device to transect the spinal cord. Before transecting the spinal cord, it is essential to understand the neurophysiology of the different anatomical structures of the nervous system. A complete transection injury event involves the dissection of both ascending and descending tracts [22]. Special
care should be taken for the pain management and urinary bladder management after a complete transection. To study specific spinal tract activity such as dorsal and dorsolateral CST, an incomplete transection (or hemisection) injury can be applied. After therapeutic interventions, this injury can also provide information regarding axonal tract regeneration. The hemisection model of injury is suitable at the level of cervical region CST which is responsible for forelimb reaching tasks. To regenerate axonal tracts for translational research after any therapeutic approach or cell transplantation, the transection model of injury can be a choice for experiments [39]. However, the transection injury model is unable to study the secondary injury mechanism.

2.1.4. Selection of an optimum injury model

Pathological and behavioral functions that alter after a SCI depend on the site and type of the injury. A correct injury model can give a proper understanding of the system and the cellular-level changes after a SCI. Furthermore, an experimental SCI model is necessary to evaluate the safety and reliability of a new therapeutic intervention before translating it to humans [40]. Every injury model has advantages and disadvantages over other models. Some selection criteria (listed in Table 1) are necessary to select the most optimum one for a specific study. For example, for translational research, contusion and compression injury models are useful as these models mimic some relevant characteristics of the lesions of traumatic injuries of the spinal cord. As they mimic real-life events, both models are also useful to study the secondary injury mechanism. Even though there are clear advantages of clinical translation of the injuries, both models fail to confirm the anatomical lesion, confidently and precisely. In contrast, transection injury model confidently cuts specific neural roots and circuits as per experimenter’s demand and thus very useful for neuroscience research [13]. However, a post-experiment histology might be required to confirm the lesion. To study a specific drug for nerve regeneration in the lesioned

| Contusion | Compression | Transection |
|-----------|-------------|-------------|
| The model is useful for neuroprotective mechanism study [114]. | Simple and reliable method [116]. | To study specific pathway function and regeneration [34]. |
| Ideal injury model for studying pathology and secondary injury mechanism [7]. | Suitable method to study secondary injury mechanism [16]. | Easy to trace the axonal tract [34]. |
| Difficult method with high variability [115]. | Useful for cell transplantation therapy | Precise control over the injury. |
| Recommended for translational research. | Difficult to control the variability between subjects. | Least variability between subjects. |
| | Suitable model for translational research [20]. | Suitable model for neuroscience research. |
spinal cord a specific nerve root lesion via incomplete transection model is suitable [41, 42]. However, for cell transplantation treatment, a compression model is recommended.

2.2. Behavioral examinations

Laboratory rats are widely used to evaluate different types of behavioral tests that involve neurological functions. After an experimental SCI in the desired location, alteration in the spinal cord and neurological function can be confirmed with behavioral assessment. Hence, behavioral examination is one of the fundamental tests after any therapeutic intervention of SCI. Different behavioral tests are carried out to estimate the pre- and post-treatment functions of various neurological tracts of the sensorimotor system [43]. According to the data collection procedure and functional organization of the nervous system these tests are categorized as: 1) endpoint measures, in which a certain behavior receives a score according to some predefined scale, e.g. number of pellets eaten in a given time; 2) kinematic measurements, which range from a qualitative description of movements to continuous quantitative measurements, e.g. Basso-Beattie-Bresnahan (BBB) scale; 3) kinetic measurements, which quantify the force produced by the limbs, e.g. forelimb grip strength tests; and 4) electrophysiological measurements, e.g. motor and somatosensory evoked potentials [43, 44]. Based on the neurological tracts used [14], these behavioral examinations can be divided into: 1) Motor test, e.g., limb grip strength test, forelimb reaching task; 2) Locomotor test, e.g. BBB scale, swim test; 3) Sensory test, e.g. Von Frey filament test, test for hot and cold sensation; and 4) Sensorimotor test, e.g ladder walking test, grid walking test, rope climbing and walking test.

2.3. Motor test

Descending tracts of motor neurons which comprises both pyramidal and extrapyramidal tracts convey information to the lower motor centers in the spinal cord that are responsible for transferring information to the muscle fiber to initiate movements of the limbs [45]. The corticospinal tract (CST) is one of the foremost descending tracts of the motor system that plays a vital role in skilled motor tasks by conveying volitional information to the limbs. The degree of CST injury is strongly co-related to the impairment of skilled motor function after a SCI [46]. After a cervical cord injury, motor control of the forelimb is profoundly affected due to the impairment of the CST. Motor functions may be restored after skilled motor learning training that assists in the reorganization of the nerve fibers [47]. To assess the motor recovery of descending tracts, several functional tests have been developed. These tests include evaluation of the motor functions of different limbs after complete and incomplete SCI. Among these tests the Irvine, Beatties and Bresnahan (IBB) scale, skilled forelimb reaching task and limb grip strength test are the most common to evaluate the
motor functions after an experimental SCI in laboratory rats. In the following sections, we discuss them in detail.

2.3.1. Skilled forelimb reaching task

The forelimb reaching task is time-consuming and needs several weeks of training to induce motor learning for forelimb food pellet retrieval. First, rats are habituated with the apparatus and special food pellet (usually sugar pellet) for one to two weeks. Food restriction is given from 3 days prior to starting the training. Training is provided for several weeks for the rats to master the reaching and grasping of the food pellets. A pellet is placed on a pit platform in front of the box slit and it must be ensured that the rat approaches the opening in a continuous manner as shown in (Fig. 2). Ten pellets for warm up and 20 pellets per task are usually used to evaluate the reaching behavior. Qualitative and quantitative assessments of a rat’s skilled reaching task are performed after 4 weeks of training (5 sessions/week) [48, 49]. For qualitative assessment, 10 components of movements (digits to the midline, digits flexed, elbow in, advance, digits extended, arpeggio, grasp, supination I and II) are rated. Animals’ reaching is rated on a 3-point scale. For normal reaching a score of “0” and for abnormal movement “1” is used. If there is no movement then “2” is given [50, 51]. For quantitative assessment, the success rate of a rat’s reaching is usually calculated. It includes the total number of pellets retrieved and eaten. The following equation is used for reaching success rate estimation:

Fig. 2. Skilled forelimb reaching task for a laboratory rat. A) A special Plexiglas chamber (40 cm × 25 cm × 30 cm) with a 1–2 cm wide opening for grasping the food pellet from a pit is normally used which is 3 cm above the base. B-E) Different stages (advance-forelimb is advanced through a slot to the platform, digit extension-forepaw is extended toward the pellet, grasp-paw grasps the pellet and supination-paw is withdrawn from the slot and a pellet is successfully taken into the mouth) of reaching and grasping of a sugar pellet by a trained Sprague Dawley rat.
Success rate = Number of pellets / Number of trials × 100

Furthermore, the amount of time required for grasping the pellets can also be measured to evaluate forelimb functional recovery [52]. Pre-injury assessment of success rates for forelimb reaching was performed for five rats. The experimental protocol was approved by the Animals Subjects Ethics Sub-committee (ASESC) of The Hong Kong Polytechnic University. Fig. 3 demonstrates the average number of food pellets grasped and consumed in each trial (20 pellets per trial) by all the five rats. After one week of apparatus habituation, the rats were trained to grasp 45 mg of dustless sugar pellets (Bio-serv®, Flemington, NJ, USA). After 42 days of training, reaching scores for each time point were compared to the untrained day 1 score. Significant differences of motor learning were found on day 22 and day 42 of the training ($P < 0.05$ and $P < 0.001$ respectively; paired t-test).

A major limitation of the skilled forelimb reaching task is that partial lesion may not impair the reaching success and qualitative scores including forepaw aiming, supination and food pellet release. However, it is also expected that complete restoration of the damaged corticospinal pyramidal tract will not restore the original behavioral condition [53].

2.3.2. Irvine, Beatties and Bresnahan (IBB) scale for forelimb function

The IBB scale is a 10-point scale to identify recovery of both proximal and distal forelimb functions after unilateral cervical SCI. Unlike the forelimb reaching task, acclimatization of rats and food deprivation is not necessary for this test [54]. First, a rat is placed in a cylindrical Plexiglas chamber for acclimation to the food. When

![Fig. 3. Success rate (Mean ± SEM) of the food-pellet reaching task of uninjured rats (n = 5). Significant differences were found after 22 days (***$P < 0.05$) and 42 days (****$P < 0.001$) of reaching and grasping training from day 1 (untrained).]
the rat is well adapted to the apparatus and food, grasping and eating behavior are recorded by a video camera. Various sizes (spherical and doughnut) of cereals are generally used to assess forelimb function. From the recorded video, elbow position (extended or flexed), supporting of the paw (contact or noncontact), position of forepaw (clubbed, extended or flexed), digit movements and grasping method are evaluated [55]. This test, however, is a qualitative assessment of the forelimb function. Visual inspections by the experimenter may affect the experimental assessment and outcomes. Sometimes, the recorded video fails to provide any information for scoring as the animal often shifts position during eating the cereals [54].

2.3.3. Limb grip strength test

A grip strength test is used to evaluate both forelimb and hindlimb functions [56]. Quantitative strength of the flexor muscles can be measured by this test. Grip strength measurement is important because it can predict any motor deficit in the limbs after a SCI. Anderson and colleagues showed that after a complete unilateral hemisection, rats lose the gripping ability permanently [57]. Acclimatization of rats to the testing apparatus is necessary before starting the test [58]. At the beginning of measurement, the rat is held by the neck or back and allowed to grab the metal grid or pull bar with either forepaw or hindpaw. The rat is then gently pulled away in horizontal plane by the tail. The sensor reads the pull strength until the rat releases the bar. Maximum grip strength is then measured from the recorded force. For each limb, several testing sessions are usually conducted to evaluate the maximum grip strength. This method is a simple procedure to evaluate the recovery over time and the response is immediate. However, after daily repeated testing, rats may hesitate to grasp if they feel an unpleasant sensation when pulled by the tail [59]. Unlike to other motor tests, in grip strength test, rats are forcibly motivated to perform the task. Hence, the measured value may differ from their natural grip strength value. The test values are also experimenter-dependent, adding more variability into the measurement. Furthermore, due to multiple measurements, the muscle may become fatigued and results may differ from each measured value.

2.4. Locomotor test

In mammals, the spinal cord contains an intrinsic locomotor mechanism that is controlled by a rhythmic central pattern generator (CPG) [60]. The location of the CPG in rats is found throughout the spinal cord, but most prominently at the lower lumbar part that rhythmically activates flexor and extensive motor pools of hindlimbs during locomotion [61]. To transmit the neuronal impulse, the dorsal root of the spinal cord carries sensory information and transmits it to the CPG for integration within the spinal cord for motor output (Windhorst 2007). During locomotion, the motor output carries information for the gait control (Grillner and Jessell...
Hence, to evaluate the recovery after SCI, locomotion tests of rats are commonly used [62]. Common locomotion tests include open-field test, swim test, BBB scale and BBB sub-scoring scale. In the following sections, we discuss these tests in detail.

### 2.4.1. BBB scale and BBB sub-scoring scale

In 1954, Tarlov and Klinger designed a qualitative scale for locomotion [63]. Due to less reliability and sensitivity, it was further modified to the Basso, Beattie and Bresnahan (BBB) scale [64] and is widely used to evaluate hindlimb locomotion function. It is a 21-point qualitative measurement developed for rats where ‘0’ refers to no observable hindlimb movement and ‘21’ refers to consistent hindlimb movement. This 21-point scale represents several stages during recovery such as movement of joints, forelimb and hindlimb coordination, trunk stability, toe clearance and tail position. From 0 to 7 is related to movement of the joints and 8 to 21 are related to planter stepping and consistent coordination of gait [64, 65].

The BBB scale has some advantages over other locomotor tests. For example, there is no training needed for this assessment. Furthermore, this test does not require any special apparatus, and thus rats can incorporate a wide range of functions [66]. BBB scale is an assessment for locomotion study which is widely used for the evaluation of interlimb coordination via reticulospinal and vestibular pathways [67]. Following a SCI, electrical and pharmacological stimulation can trigger the rhythmic circuit to generate locomotion type movements [68]. For studies of reticulospinal regeneration after a thoracic cord injury, the BBB scale is a good method to evaluate the assessment. Furthermore, rhythmic activity can give information on excitatory interneurons of the spinal cord. In the case of therapeutic treatment, the scale does not follow the order of the BBB scale [69]. Furthermore, for non-contusive models of SCI, many functional state conditions cannot represent the BBB scale [70].

To further solve the sensitivity of the BBB scale and to evaluate the higher motor function, a 7-point sub-scoring system was developed by Lankhorst and colleagues. In this sub-scoring scale, all the behaviors are cumulated independently and then added together to yield a single score [71]. The major limitation of the BBB sub-score is that it is only valid for mildly injured animals [72].

### 2.4.2. Open field test

The open field test is also the simplest method to evaluate locomotor behavior and anxiety in rats after a SCI. In 1934, Calvin Hall developed a test where defecation was used as a measure of anxiety [73]. It is also used to evaluate the efficacy of therapeutic drugs on anxiety-like behaviors in rodents [74]. To evaluate the exploratory behavior deficit after a SCI, open field test of rat is also used [75].
closely controlled conditions a square open field arena (60 × 60 cm), made of black Plexiglas is required to assess behavior. The bottom of the area is subdivided into 12 × 12 cm squares. A rat is placed at the center of the area and the number of squares crossed is counted and recorded [76]. Additionally, the open field test is also used to measure the locomotor recovery of hindlimbs [77]. Another modification to this test named open field motor test (OFT) was also developed for motor evaluation [78, 79].

2.4.3. Swim test

The swim test is a standardized natural evaluation method of locomotor recovery after SCI [80, 81]. The method is also suitable for measuring the spasticity after a con- tusive SCI [82]. Rebecca and colleagues developed the Louisville Swimming Scale (LSS) which mainly evaluates forelimb and hindlimb movements and body position. It cannot, however, uniformly evaluate mild to moderate SCI in rats [83]. To combat this problem, in 2015, Xu and colleagues developed a swimming scale to assess the locomotor recovery after a SCI in rats [72].

For the swim test, typically a Plexiglas swim tank (150 cm × 15 cm × 30 cm) is used [72]. The tank is usually filled with tap water (28–32 °C) to a depth of 20 cm. To observe rats during swimming, a mirror is placed underneath the tank at a 45-degree angle. A total of five trials per rat are used to evaluate the swimming behavior. Forelimb and hindpaw movements, and the distance between hindpaw and tail are used to analyze the swimming behavior. All the parameters are measured on an 8-point scale as described previously [84]. An important limitation of the test is that it cannot be performed in complete SCI animals. Moreover, the rats need extensive training to obtain effective results; otherwise it may induce depression behavior [85].

2.5. Sensory test

Sensory feedback such as afferent proprioception plays a significant role in the control of motor functions. Hence, sensory tests are used to evaluate the hyperactivity of rats in the presence of heat [86], cold [87] or pain [88]. After an experimental SCI and therapeutic interventions, sensitivity tests are useful to evaluate the sensory pathways of the experimental rats [89]. By selecting a correct hindlimb sensory test for these SCI rats, a sensory assessment can be carried out. In the following sections, we further discuss some of these sensory tests in detail.

2.5.1. Tactile sensory test or neuropathic pain test

To test hindlimb sensation, the tactile sensory test is used by determining whether the rat can stand or not after a SCI [90]. A group of special filaments (Von Frey
Filament) are used to evaluate the improvement of neuropathic pain after therapeutic interventions in SCI rats [91]. Two validated testing methods - up/down plantar Von Frey (VF) test, and dorsal Von Frey Hair (VFH) monofilaments - are used for the evaluation [90]. Before the evaluation, rats are habituated with the cage by providing cereal grain so that they can seek food for a reward. The up/down plantar VF test is calibrated by VF microfilaments ranging from 0.4–74 gm to elicit paw withdrawal by the stimulus. For a successful test, at least 10 sequential trials are required for each paw, and about 3–5 min of resting period must be maintained between two trials. In the dorsal VF test, first the rat is wrapped in a towel so that the hindquarters are exposed. The stimulus is then given to the dorsal surface in between the first and second metatarsal by the VFH filaments. Withdrawal of the paw indicates that the spinal reflex is active. Hair value for left and right paw are recorded and withdrawal responses are marked as positive and negative. Both of these forces produced by the Von Frey Hairs are then averaged to yield a single score [92]. The VFH test has high sensitivity to assess the tactile sensation of the hindpaws, and it can also give information on neuropathic pain in chronic conditions. A disadvantage of this test is, however, that it requires a very long time of repetitive measurements which may lead to sensitization or learnt responses [93].

### 2.5.2. Test for hot sensation

This is a modification of the tail flick test where a heating beam is placed under the tail for a predefined time to assess the thermal sensation [94]. Later, a preheated (50 °C) plate is used for this test to evaluate the sensory stimulation of hind paws after a thoracic injury. A rat is placed on this hot plate for a maximum of 60 seconds. If the rat shows any sign of licking its hind paws within this time, the rat is considered as positive. If the rat does not show any sign of licking, it is considered as a negative and must be removed from the hot plate after 60 seconds [95]. A major limitation of the test is that the rat may vocalize or be injured due to the heat or burn its paw skin. The problem can be overcome by wrapping the rat in a towel and placing only the plantar on the metal surface [96].

### 2.5.3. Test for cold sensation

Unlike the hot sensation test, the cold sensation test is also useful for sensory nociceptor testing. Ethyl chloride is mainly used for testing the cold sensation of rats. First, the area of interest of the skin of a rat is shaved. Then ethyl chloride is sprayed onto it, and the response is recorded and graded based on the following scale: 0 = no response, 1 = localized response, 2 = transient vocalization, and 3 = sustained vocalization [88]. Other chemicals such as acetone can also be used for cold sensation evaluation. Application of 100 μl of acetone onto the plantar hind paw was applied to test the cold response in rats [87]. In addition, this sensory test can
evaluate only unilateral pain measurement. A complete injured animal model is not suitable due to the variability during measurements.

2.6. Sensorimotor test

Unilateral or bilateral damage to the spinal cord is responsible for chronic deficits of somatosensory functions [97] and asymmetry of limb movements and inter-limb coordination [98]. The somatosensory system also incorporates the afferent, efferent and central integration pathways of the spinal cord to maintain homeostasis during joint movements. To evaluate this sensory and motor integration after a SCI and treatment, some sensorimotor tests are commonly used in rats. In the following sections, we discuss these tests in further detail.

2.6.1. Horizontal ladder walking test

To evaluate forelimb and hindlimb stepping and coordination, a horizontal ladder walking test is frequently used [99]. Special ladder apparatus and video recording are needed for quantitative and qualitative analysis of walking. Two horizontal Plexiglas walls (100 cm × 20 cm each) with 50 small holes inserted with plane metal rungs are generally used to make the horizontal ladder (Fig. 4). The rungs are placed in a regular pattern at about 2 cm intervals. Rats are allowed to cross the horizontal ladder from a new cage to the native cage. A reward is given after each successful cross across the ladder.

After several days of training, different templates of irregular rung patterns are used to normalize the difficulty and enhance comparability during learning [70]. The recorded videos of the rat’s footsteps on the rungs are used for analyses. A 7-category foot fault scoring scale (0-total miss, 1-deep slip, 2-slight slip, 3-replacement, etc.) is used.

**Fig. 4.** Ladder rung test for rats. A) Two horizontal Plexiglas walls with 50 small inserted metal rungs; B) The rat completely misses the metal rung when walking along the ladder; C) Rat lifts its limb and replaces it on another metal rung.
4-correction, 5-partial placement and 6-correct placement) is used for the quantitative measurement of foot paw placement on the rungs.

2.6.2. Grid walking test

Grid walking test is useful to evaluate and predict the efferent and afferent sensorimotor integration [100]. There are several versions of this test such as horizontal grid walking [44], ladder beam [101], inclined grid climbing [102] etc. These tests are useful for measuring sensorimotor deficit and recovery after any experimental injury and treatment.

The grid walking test is an another method to assess the sensorimotor deficits and motor incoordination in rats after a SCI [100]. The test is useful for animals with chronic neurological deficits. Because foot faults are quantified during the test, it is also termed as the foot fault test [103]. A rat is placed on a metal grid and allowed to cross a 1–1.2 m long runway with irregular gaps. A fence is used to restrict the roaming of the rat. The rat is required to put its limbs accurately on certain places. When the paw of a rat falls down, it is considered as a foot fault. Errors are counted during the walking on the grid for 1 minute. A 3-point score is given to evaluate the task. In complete transected or severely contused animal models, a major limitation is that the differentiation of descending spinal influences may be difficult; however, this measure was found to be a sensitivity indicator for small residual neurological deficits [66, 78].

For inclined ladder climbing, a ladder made up of plastic frame (1280 mm × 190 mm) with 19 round rungs (6 mm in diameter) is placed in an inclined position (55°). Training of animal is essential for this test. After successful training, rat is placed at the bottom of the ladder and climbing video is recorded. From the video, hindpaw placings are evaluated and correct ladder steps are noted. The test can provide an evaluation of motor behavior after severe paralysis. However, the sensory feedback cannot properly evaluated by this test. Furthermore, marginal improvements of hindlimb stepping also cannot be evaluated properly; although, for horizontal ladder walking, the hindlimb stepping can be evaluated properly.

2.6.3. Rope climbing and walking test

Pre-training of rats is needed for the rope climbing test. During this test, rats are required to climb 160 cm and 1.25-in. wide vertical rope four times. Videos are recorded using a camera to evaluate the climbing. From the video, the total number of foot slips per rope climbing are evaluated for 42 days after the injury [104]. For the horizontal rope walking test, rats are trained to cross an 83 cm horizontal rope as described previously [105]. Rats are then tested for five consecutive trials. Quantitative assessments of the rats are performed based on slips and falls from...
the rope during each trial. The total number of errors per 10 steps is analyzed to evaluate the locomotion on a 4-point rating (0- normal locomotion, 1- close to normal locomotion, 2- moderate or unable to achieve continuous placement, 3- frequent falls and slips).

The sensorimotor test is useful to assess gait during walking. To find out the coordination of both sensory and motor functions, ladder walking tests are performed before and after a SCI. With special apparatus, bipedal movement capacity can also be carried out during locomotion [106]. Nonetheless both quantitative and qualitative assessments are essential to validate the study after an experimental SCI.

2.6.4. Selection of an appropriate behavioral test

An appropriate behavioral test should be chosen to understand the effects of new therapeutic interventions on experimental animals. Limitations of the behavior tests should be considered before proceeding with the test. For specific tract recovery, a behavior test should be chosen very wisely. For example, to evaluate the motor recovery, especially after CST injury, skilled reaching task in rats is useful. Similarly, for drug and cell transplantation treatments for sensorimotor recovery, different grid walking tests can be considered. Behavioral assessment should also be chosen on the basis of reliability, validation and sensitivity [64, 107, 108]. The overall criteria for selecting a particular behavioral test are summarized in Table 2.

A healthy nervous system utilizes all the functions to accomplish one particular task, it is expected that not all the functions are dominant to that task. For example, while reaching task is dominant with motor function and walking task is dominant with locomotor function; for natural reaching and walking, along with motor and locomotor functions, both sensory and sensorimotor functions are also essential. However, one or two appropriate tasks are sufficient to test the recovery of one particular function.

**Table 2. Summary of broad four types of behavioral tests and their selection criteria.**

| Motor test | Locomotor test | Sensory test | Sensorimotor test |
|------------|----------------|--------------|-------------------|
| - To study volitional movements including arm and digit movements [54]. | - To study the central pattern generator (CPG) mechanism [117]. | - To examine sensory integrity and improvement after any neurological injury [90]. | - To examine somatosensory and sensorimotor integration [119]. |
| - Qualitative and quantitative assessment of forelimb function [48]. | - To study flexor and extensor neural network organization [118]. | - To evaluate the afferent tract function [89]. | - To assess gait dynamics [106]. |
| - To test different neuronal tract functions [50]. | | | - To evaluate the coordination between left and right limb movements [99]. |
3. Conclusions

Laboratory rat with experimental SCI is a well-accepted animal model, frequently used worldwide for SCI-related translational research. Several surgical methods such as compression, contusion and transection models have been developed and tested on laboratory rats. Selection of a correct injury model is also essential to understand similar clinical conditions in patients. Plasticity of neural tract after SCI is very important and is a major topic in clinical research. However, there are still some questions needed to be answered. How the behavior or activity modulates the neural circuits and the synaptic strengths? What is the mechanism underlying this neuroplasticity? Each behavioral test, however, can answer only one or two specific questions. Hence, some new tests should be included into the experimental toolkits. For example, for studying bipedal locomotion, rats can be trained on a body-weight support system for locomotor training [109]. Recent studies have demonstrated successful translation of the research results to early-stage clinical interventions [110, 111]. For rat’s somatosensory test, tickling test, a new method of eliciting ultrasonic vocalization by gently caressing to the trunk area, can be used to test the integrity of the neuronal tract for conveying sensory information [112]. For successful clinical tests of new interventions for SCI, an appropriate experimental plan is also needed. Nonetheless, very high ethical standards should be maintained for animal welfare and for the acceptance of the scientific outcomes.

Declarations

Author contribution statement

All authors listed have significantly contributed to the development and the writing of this article.

Funding statement

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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