Review Article

Role of Acupuncture in the Management of Severe Acquired Brain Injuries (sABIs)

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Acupuncture therapy has been used to treat several disorders in Asian countries and its use is increasing in Western countries as well. Current literature assessed the safety and efficacy of acupuncture in the acute management and rehabilitation of patients with neurologic disorders. In this paper, the role of acupuncture in the treatment of acute severe acquired brain injuries is described, acting on neuroinflammation, intracranial oedema, oxidative stress, and neuronal regeneration. Moreover, beneficial effects of acupuncture on subacute phase and chronic outcomes have been reported in controlling the imbalance of IGF-1 hormone and in decreasing spasticity, pain, and the incidence of neurovegetative crisis. Moreover, acupuncture may have a positive action on the arousal recovery. Further work is needed to understand the effects of specific acupoints on the brain. Allegedly concurrent neurophysiological measurements (e.g., EEG) may help in studying acupuncture-related changes in central nervous system activity and determining its potential as an add-on rehabilitative treatment for patients with consciousness disorders.

1. Introduction

Severe acquired brain injuries (sABIs) include a variety of acute brain lesions characterized by the occurrence of variably prolonged coma (24 hours), and simultaneous motor, sensory, cognitive, and/or behavioural impairment that causes a certain degree of disability. Congenital, perinatal onset, or degenerative-progressive brain injuries are excluded from this definition.

A common consequence of sABIs is disorders of consciousness (DOCs), a prolonged cognitive impairment including the loss of awareness of oneself and environment. DOCs represent one of the greatest challenges that modern medicine faces today, with a huge burden of care for families and facilities. On the basis of the current taxonomy of DOCs, a state of altered consciousness can be categorized into coma, vegetative state (VS), also referred to as unresponsive wakefulness syndrome (UWS), and minimally conscious state (MCS) [1]. The most common cause of sABI is traumatic brain injury (TBI), a major source of death and disability worldwide. Two further causes of sABI are anoxic encephalopathy (AE), usually due to cardiocirculatory arrest (secondary to extensive myocardial injury and/or malignant arrhythmias), and ischemic or haemorrhagic stroke. These conditions mostly affect subjects from the fifth decade onwards and represent about 40% of the sABIs. The main involvement of neurons leads to a worse prognosis than TBI. In AE, the neurons disruptions, with a low regenerative potential, cause a high risk of irreversibility of the consciousness disorder. Moreover, the time interval within which a recover is reasonable, up to 12 months from the event for TBI, is unlikely more than 3-6 months from the event for anoxic or vascular brain injuries. Late recoveries are possible, but rare. Further nontraumatic sABI arises from brain tumors, infections, and toxic-metabolic encephalopathy [2].

Despite the fact that steady progress has been made toward prolonging patients survival and several
Table 1: Physiopathology of DOCs.

<table>
<thead>
<tr>
<th>Primary brain injuries</th>
<th>secondary brain injuries</th>
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<tbody>
<tr>
<td><strong>Focal</strong></td>
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<tr>
<td>(i) contusion</td>
<td>(i) ionic imbalance (due to hypoxia and hypotension)</td>
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<tr>
<td>(ii) laceration</td>
<td>(ii) glutamate excitotoxicity</td>
</tr>
<tr>
<td>(iii) hemorrhage</td>
<td>(iii) oxidative stress (generated by mitochondrial dysfunction)</td>
</tr>
<tr>
<td><strong>Diffuse</strong></td>
<td></td>
</tr>
<tr>
<td>(i) concussion</td>
<td>(iv) ischemic injury</td>
</tr>
<tr>
<td>(ii) diffuse axonal injury</td>
<td>(v) edema formation</td>
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<tr>
<td></td>
<td>(vi) intracranial hypertension</td>
</tr>
<tr>
<td></td>
<td>(vii) neuroinflammation</td>
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<tr>
<td></td>
<td>(viii) blood-brain barrier disruption</td>
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</table>

pharmacologic and neuromodulating strategies have been proposed, results on functional recovery of DOCs are still scarce.

2. Physiopathology of DOCs

The immediate effect of the impact forces on intracranial tissues, i.e., the primary brain injury, can be focal or diffuse. Focal injuries, such as contusion, laceration, and haemorrhage, are early detectable upon imaging and their consequences depend on their location and severity. Diffuse brain injuries, like concussion or diffuse axonal injury, require magnetic resonance to be detected [3]. However, what is susceptible of treatment is secondary brain injury (SBI), the cascade of biochemical and cellular events developing minutes to months after the insult: ionic imbalance (due to hypoxia and hypotension), glutamate excitotoxicity, oxidative stress generated by mitochondrial dysfunction, ischemic injury, edema formation, intracranial hypertension, neuroinflammation (by both systemic and central neuron system immunoactivation), and blood-brain barrier (BEE) disruption [3, 4], as schematized in Table 1.

Several secondary processes are based on intracellular calcium overload, due to excitatory amino acids and inflammatory cytokines released. In absence of oxygen, glucose enters the glycolytic pathway, where it is converted to NADH and pyruvate; the latter becomes lactic acid just producing 2 ATP molecules, while Krebs cycle is precluded. Thus, a downregulation of ATP-dependent Na⁺/K⁺-ATPase pump leads to sodium overload, which acting on Na⁺/Ca²⁺ exchange pump provides calcium retention.

Analogously, the huge release of glutamate following a head trauma, in hypoxic conditions, is not counterbalanced by astrocyte reuptake of glutamate, which acts on N-methyl-d-aspartate (NMDA) receptor, thus increasing calcium influx. Calcium is therefore sequestered by mitochondria causing their dysfunction, lysis, and release of byproducts [3]. Moreover, calcium stimulates neuronal and endothelial cells overproduction of nitric oxide (NO), which can be associated with oxidative stress. A further effect of calcium overload is mitochondrial-mediated apoptosis following the release of cytochrome c.

In response to primary brain injury, an inflammatory response is provided by microglia and astrocytes, which attract leukocytes via cytokines, chemokines, and NO. The disruption of the BEE often leads to cerebral oedema and intracranial hypertension [3].

3. Reactions to Brain Injury

A series of mechanisms are activated at neuronal (neogenesis, synaptogenesis, and dendritic remodelling) glial, and vascular level, referred to as neuroplasticity [5]. These events are promoted by increased expression of growth factors involved in brain development, such as nerve growth factor (NGF), neurotrophin 4/5, basic fibroblast growth factor, and brain derived neurotrophic factor (BDNF); an increasing importance has also emerged for insulin-like growth factor 1 (IGF-1), regulating metabolic function, oligodendrocyte proliferation, and survival, angiogenesis, myelinisation, and neurite outgrowth [5] and receptors for IGF-1, virtually ubiquitous, are mainly expressed by mesenchymal origin-derived cells in the hippocampus, parahippocampus, amygdala, cerebellum, and cortex [6].

4. Available Strategies to Manage sABIs

New insights into the pathophysiology of sABIs initiated new therapeutic approaches (neuroprotective strategies) aimed at interrupting secondary brain injury development and promoting mechanisms of arousal (see Table 2). Among the neuroprotective strategies, mild hypothermia (which decreases cerebral metabolic demand, excitatory neurotransmitter release, inflammation, and oedema), hyperosmolar therapies (also known for their immune-modulating property), statin (reducing oxidative stress and inflammation), and cyclosporin A (ameliorating mitochondrial function) have been proposed [3]. For arousal recovery, dopaminergic, serotoninergic and GABAergic drugs have been explored, with occasional results. Several nonpharmacological strategies have been utilized. Early verticalization has been shown to increase the spectral power of the EEG higher frequencies and the subject’s arousal [7, 8], probably enhancing vestibular afference on locus coeruleus, raphe, and thalamic intralaminar nuclei. Enriched environment, equipped with emotional stimuli and biographically meaningful objects, showed greater range of behavioural responses [9]. Finally neuromodulatory techniques, including deep brain stimulation (DBS), transcranial direct current stimulation (TDCS),...
Table 2: Available strategies to manage sABIs and their mechanisms of action.

<table>
<thead>
<tr>
<th>NEUROPROTECTIVE STRATEGIES</th>
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<tbody>
<tr>
<td>Mild hypothermia</td>
<td>decrease of cerebral metabolic demand, excitatory neurotransmitter release, inflammation and edema</td>
<td></td>
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<tr>
<td>hyperosmolar therapies</td>
<td>immune-modulating effects</td>
<td></td>
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<tr>
<td>statins</td>
<td>reducing oxidative stress and inflammation</td>
<td></td>
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<tr>
<td>cyclosporin A</td>
<td>ameliorating mitochondrial function</td>
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<tr>
<th>NEUROMODULATIVE STRATEGIES</th>
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<tbody>
<tr>
<td>Pharmacologic</td>
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<tr>
<td>(i) dopaminergics</td>
<td>neurotransmitter modulation</td>
<td></td>
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<tr>
<td>(ii) serotoninergics</td>
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<td></td>
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<tr>
<td>(iii) gabaergics</td>
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<tr>
<td>Non Pharmacologic</td>
<td>potential enhancement of the vestibular effects on locus coeruleus, raphe and thalamic intralaminar nuclei</td>
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<tr>
<td>postural verticalization</td>
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<tr>
<td>Neuromodulative Techniques:</td>
<td>increasing metabolism in the forebrain, thalamus and reticular formation. modulating neuronal networks and ANS</td>
<td></td>
</tr>
<tr>
<td>(i) deep brain stimulation (DBS)</td>
<td></td>
<td></td>
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<tr>
<td>(ii) transcranial direct current stimulation (TDCS)</td>
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<td></td>
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<tr>
<td>(iii) transcranial magnetic stimulation (TMS)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(iv) spinal cord stimulation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(v) median nerve stimulation</td>
<td></td>
<td></td>
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<tr>
<td>(vi) vagus nerve stimulation</td>
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</tbody>
</table>

Transcranial magnetic stimulation (TMS), spinal cord stimulation, median nerve stimulation, or vagus nerve stimulation, have been proposed.

5. A Complementary Approach for the sABIs’ Management Deriving from Ancient Chinese Medicine: Acupuncture

The aim of this review is to explore the potential role of acupuncture (1) in the acute treatment of sABIs, in containing secondary brain injury by acting on different pathophysiological mechanisms, and (2) in the subacute/chronic management of sABI outcomes, through a limitation of spasticity, pain, dysautonomia, and a possible action on the arousal recovery.

Acupuncture is a traditional medicine traced back to over 3000 years ago in China [10], consisting in inserting needles into specific points of the patient’s body ("acupoints") chosen on the basis of the Meridian theory of Traditional Chinese Medicine (TCM) [11]. It has been used in the treatment of stroke and its consequences for over 2,000 years in China.

Under the influence of neuroanatomy, neurophysiology, and bioholographic principle of modern medicine, in the early 1970s, scalp acupuncture, one of the several specialized acupuncture techniques in which a filiform needle is used to penetrate specific stimulation areas on the scalp mainly for the treatment of brain diseases, was set up and separated from traditional acupuncture system [12].

Scalp acupuncture has been reported to (1) improve cerebral blood circulation, promoting regional energy metabolism; (2) upregulate expression of glial cell-line derived neurotrophic factor (GDNF), possibly promoting proliferation and differentiation of neural stem cells in the focal cerebral cortex and hippocampus; (3) reduce contents of excitatory amino acid and increase level of GABA, thus lowering neurogenic toxicity; (4) ease cerebral vascular immunoinflammatory reactions; (5) regulate blood lipid metabolism to resist cerebral free radical damage; and (6) inhibit cerebral cortical apoptosis [13].

5.1. Acupuncture and Inflammation in DOCs. Few studies considered the diachronic assessment of inflammatory markers after a sABI in order to monitor the local and systemic stress response in DOCs. Amico and colleagues evaluated the subacute phase of DOCs in five patients in vegetative state (VS) and one in minimal consciousness state (MCS), by clinical assessment and biochemical analyses [14]. A positive correlation was found between the serum levels of osteopontin (OPN), a cytokine involved both in neuroinflammation and neurorepair, and prognosis, with the lowest level detected in the patient who then emerged from MCS, the highest in the one who then died. Moreover, the lymphocyte subset presented a general increase of CD4+/CD3+ ratio, with a suspect unbalance of CD4+ toward Th2; prolactin resulted to be the best endocrinological marker of sABI [14].

Applied at Baihui (GV20) and left Zusanli (ST36), acupuncture significantly reduced the infiltration of inflammatory cells and the expression of the proinflammatory enzyme MMP2 in cerebral ischemia/reperfusion injury (CIRI) model rats. In particular they attenuated the expression of the water channel proteins P4 and AQP9 in the ischemic brain, leading to the mitigation of inflammation-related brain edema. Consistent with the smaller observed infarct size, acupuncture and EA both promoted significant improvements in the Modified Neurological Severity Score (mNSS) in CIRI model rats, indicative of enhanced neurological function [15].

Moreover, acupuncture successfully downregulates tumor necrosis factor alpha (TNF-α), which results in
5.2. Acupuncture and Redox Equilibrium. Oxidative stress, the imbalance between the production of reactive oxygen and nitrogen species (ROS/RNS) and the endogenous antioxidant system, causing a cascade of chain reactions resulting in cellular damage, is a critical feature in the pathological process of various diseases [17].

Recently, a large body of evidence demonstrated that acupuncture has antioxidative effect in various conditions [18–20], although the exact mechanism, especially the influence of acupuncture on signalling pathways, remains unclear. Through redox system, antioxidant system, anti-inflammatory system, and nervous system, acupuncture could make the oxidative damage and the antioxidant defence remain in a relatively constant redox state. However, the recent acupuncture researches about oxidative stress are sporadic and preliminary [21].

5.3. Acupuncture and Intracellular Calcium. The insertion of the needle represents an effective mechanical stimulus, leading to tissue displacement and to intracellular calcium increase and signalling. The modulation of calcium channels seems to be the primary mechanism for endorphin secretion and release from immunocytes and for the inhibitory effects of opioids on peripheral neurons [22]. Thus, calcium ion, whose increase is so crucial in the pathogenesis of SBI, may be taken as the carrier of the biological modulation system provided by acupuncture, where the mechanical wave onsets an acoustic shear wave, and this drives to calcium signalling [22].

5.4. Acupuncture and Neuron Regeneration. In the brain of human adults, neural stem cells (NSCs) have been demonstrated in the pallium, subependymal region, hippocampus, and corpus striatum, which have the ability of self-duplication, self-regeneration, and differentiation into neurons and glial cells. During cerebral ischemia reperfusion, astrocytes play a crucial role in limiting neuronal lesion, as they release epoxyeicosanoic acids in order to enlarge brain vessels, release Nerve Growth Factors to make neurons survive and axons grow, produce neurotransmitters, metabolize toxic molecules, and have also the potentiality to become NSCs [23, 24].

The astrocyte activation and proliferation marker is Glial Fibrillary Acidic Protein (GFAP), while Neuron Specific Enolase (NSE) is one of the neurons' markers. Acupuncture on the conception (CV) and governor vessels (GV) has been shown to inhibit excessive proliferation of astrocytes and promote NSCs differentiation in the ischemic brain [25, 26]. In particular, needling acupoints GV20 and GV26 could downregulate the number of GFAP+ cells, while increasing the GFAP/NSE double–labelled cells in the hippocampal dentate gyrus [25, 26]. Another study, which employed a rat TBI model, proved that during the early post-TBI stage, acupuncture (GV20, GV26, GV15, GV16, and LI4) can promote the proliferation and differentiation of NSCs and glial cells, which is crucial to control neuronal necrosis; in the late phase, it can inhibit glial proliferation and differentiation, driving to neuron and oligodendrocytes regeneration and tissue repair [27].

Moreover, needling CV24, CV4, and CV3 has been shown to upregulate the expression of basic FGF, EGF, and NGF after cerebral ischemia reperfusion, activating nerve repair and proliferation of neuronal precursors [28, 29].

As regards human studies, recent evaluations used single-photon emission computed tomography (SPECT) and T2-weighted imaging (T2WI) [30], or functional magnetic resonance imaging (fMRI) [31]. Shen and colleagues compared serial diffusion tensor imaging (DTI), fluid-attenuated inversion recovery (FLAIR), and T2WI performed in 20 patients with recent cerebral infarction in the basal ganglia, randomly divided into an acupuncture group and a control group [32]. The apparent diffusion coefficient (ADC) of infarction lesions, decreased at stroke onset, was showed significantly elevated after the acute stage, while the ADC of the bilateral cerebral peduncle was reduced on the infarction side. Fractional anisotropy (FA) values of abnormal signals on DTI in the infarction areas and cerebral peduncles underwent a significant reduction from stroke onset to the chronic stage. Interestingly, a significant difference in ADC and FA values between the two groups was observed, with a higher FA value in the acupuncture group than the control group, thus suggesting the effectiveness of acupuncture for protecting neurons by postponing Wallerian degeneration of brain infarction, and facilitating recovery [32].

5.5. Acupuncture and Arousal. The pathology of disorders of consciousness can be represented by (A) damage of Reticular Ascending System (B) large-scale damage to cerebral cortex, (C) injury to links (e.g., thalamus) between cerebral cortex and brain stem, and (D) injury to connections (e.g., corpus callosum) within the cerebral cortex, i.e., severe diffuse axonal injury (DAI).

The production of inhibitors (including GABA) induced by brain injury generates a response resembling automatic shutdown, probably aimed at conserving energy and promoting cell survival, but causing a comatose state [33]. Therefore, any treatment affecting the reticular activating system may be worth trying, and, among the possible treatments, acupuncture has the highest potential [34].

Recent studies on resting state (RS) in DOC, by using functional magnetic resonance imaging (fMRI), showed that functional connectivity is severely impaired above all in the Default Mode Network (DMN). In the vegetative and minimally conscious state, DMN integrity seems to correlate with the level of remaining consciousness.

The DMN is among the most robust networks found with resting state fMRI and encompasses the posterior cingulate cortex (PCC)/precuneus, medialfrontal/anterior cingulate cortex, and temporoparietal cortex [35]. Activity in the
DMN diminishes when the brain is involved in attention-demanding cognitive tasks [36] and returns to its prominent presence when no such task is being performed.

The DMN seems to be of particular interest, as its connectivity has been shown to decrease during loss of consciousness, and PET studies have shown an increase in neuronal activity in DMN regions (especially in the PCC/precuneus) upon recovery from VS [37]. Indeed, Vanhaudenhuyse [38] and Fernández-Espejo [39] et al. observed a correlation between the DMN integrity and the level of consciousness in noncommunicative brain-damaged patients.

Imaging evidence has been provided to support that electroacupuncture at GV20, employed to treat major depressive disorders, may modulate the Default Mode Network (DMN), the cerebral functional network encompassing the posterior and anterior cortical midline structures, which is considered to be involved in stimulus-independent thought, mind-wandering, and self-consciousness. EA at GV20 would increase functional connectivity (FC) between the precuneus/posterior cingulate cortex (PC/PCC) and bilateral anterior cingulate cortex (ACC) and reduce FC between the PC/PCC and left middle frontal cortex, left angular gyrus, and bilateral hippocampus/parahippocampus [40]. These findings are of particular importance when considering DOCs, where resting state network activity reveals reduced interhemispheric connectivity and correlates with levels of consciousness.

The acupoint Shuigou (GV26), placed at the junction of the upper one-third and lower two-thirds of the philtrum midline, also has been described as promoting the function of GV meridian, closely related to brain function, decreasing cognitive impairment, and promoting neurogenesis in the APP/PS1 transgenic mice [41].

Interestingly, enhanced bodily awareness can be triggered by genuine acupuncture at PC6 and HT7 acupoints, which were exhibited to activate the salience network (insula, ACC, secondary somatosensory cortex, and superior parietal cortex) and deactivate the DMN (medial prefrontal cortex, PCC, inferior parietal cortex, and parahippocampus) [42].

Combined with western medicine, electroacupuncture therapy at Baihui (GV20), Shuigou (GV26), and Yongquan (KII), resulted effective in improving consciousness recovery of patients in coma due to TBI, both reducing awake time and increasing awake rate, compared to a control group receiving only western treatments [43].

5.5.1. Autonomic Dysfunction in sABI. Autonomic nervous system (ANS) deregulation and/or dysautonomia is another severe consequence of brain injury, not well cleared. Dysautonomia affects in particular ninety percent of TBI patients during the first week, leading to sleep and heart rhythm disorders, and increasing specific biomarkers of neural damage [6]. Clinically, patients suffering from DOC can show the so called “paroxysmal sympathetic hyperactivity” (PSH), episodic sudden increase in vital signs, particularly heart rate, blood pressure, respiratory rate, and temperature, with possible diaphoresis (i.e., excessive sweating) and abnormal, unintentional movements, spontaneously or in response to external painful stimuli [44, 45]. A growing body of evidence suggests that ANS may mediate large-scale brain activation, in an extreme attempt to preserve body system homeostasis and regain consciousness [46]. The primary and secondary brain lesions have the potential to compromise both cortical and subcortical control mechanisms of the ANS.

Most often, TBI leads to sustained sympathetic activation, contributing to the high morbidity, with oxidative stress in the ANS and activated hypothalamic-pituitary-adrenal axis and hypothalamic-sympathoadrenal medullary axis [4].

5.6. Acupuncture and ANS Modulation. An increasing clinical evidence demonstrates that acupuncture is helpful in treating ANS dysfunctions, such as nausea and vomiting [47]. For example, acupuncture stimulation has been shown to change the sympathovagal balance toward vagal predominance [48, 49].

Abnormalities in the ANS, such as sympathetic overactivation and/or parasympathetic hypoactivation, may generate and sustain chronic pain [50, 51].

Acupuncture at certain points could reduce sympathetic nervous system activity associated with pain [52] or during mental stress in patients with heart failure [53]. However, the neurobiological basis of these effects is not yet clear [54]. In order to explore the regulatory effect on ANS by acupuncture, Sakatani and colleagues monitored heart rate by placing photoelectrical sensor on the first finger of eighteen healthy male adults, and thus low frequency (LF) amplitude (0.04–0.15 Hz) and high frequency (HF) amplitude (0.15–0.4 Hz) were calculated by power spectral analysis [54]. Real acupuncture performed at point Large Intestine 4 (LI4) of the right hand (r-LI4) was shown to determine significant decreases of HR and LF/HF and a significant increase of HF, indicating a parasympathetic activation as well as sympathetic depression [55].

Moreover, vagus nerve stimulation (VNS) increases metabolism in the forebrain, thalamus, and reticular formation [56]. It also enhances neuronal firing in the locus coeruleus, which leads to massive release of norepinephrine in the thalamus and hippocampus, a noradrenergic pathway essential for arousal, alertness, and the fight-or-flight response [57]. Recently, based on this rationale, Transcutaneous Auricular VNS (tVNS), a noninvasive stimulation developed for treating epilepsy and depression without the surgery-related risks [58, 59], was firstly employed by Yu-tian Yu and colleagues on a patient in vegetative state [60]. A further case with implanted VNS recovered behavioural responsiveness and enhanced brain connectivity patterns [61].

5.7. Acupuncture against Neuroendocrine Dysfunction. An imbalance of the pituitary and hypothalamus hormones and their axes is often associated with sABI, due to compression, edema, skull base fracture, haemorrhage, intracranial hypertension, or hypoxia. In the acute phase of sABI, low IGF-1 with elevated GH levels have been detected, with increasing IGF-1 and normalizing GH concentrations in the following weeks. The IGF-1 upregulation and the disruption of BEE that persists until 7 days after injury, allowing a wide level of hormone to reach the surviving neurons, probably play a
role in promoting neurite overgrowth, inducing progenitor cell differentiation and inhibiting neuron apoptosis [6].

In rat model of renal failure (RF)-induced hypertension, stimulation with acupuncture, and most significantly with EA, at ST36 and KI3, not only attenuated glomerulosclerosis and tubulointerstitial fibrosis, but corrected the decreases in RF-induced IGF-1 mRNA and protein levels, thus counteracting oxidative stress [62]. These findings may suggest the ability of acupuncture in restoring IGF-1 function in any situations where its levels are reduced, including sABI, although no studies have been conducted on this purpose.

5.7.1. Pain and DOC. The experience of pain in disorders of consciousness is still debated. Neuroimaging studies, using functional magnetic resonance imaging (fMRI) [63, 64], Positron Emission Tomography (PET) [65], multichannel electroencephalography (EEG), and laser-evoked potentials [66], suggest that the perception of pain increases with the level of consciousness.

VS and MCS are by definition incompatible with a reliable and consistent ability to communicate about pain experiences, while the nature of these conditions is characterized by various factors that can give rise to pain (e.g., spasticity, contractures, etc.) [67].

5.8. Acupuncture and Pain Relief. Many different mechanisms may explain the analgesic effect of acupuncture. Among these is the gate control theory of pain proposed by Melzack and Wall in 1965 [68]. Specific nerve fibers would transmit pain to the spinal cord, while other nerve fibers inhibit pain transmission. Both groups of fibers met at the substantia gelatinosa in the spinal cord, where pain and pain inhibitory stimuli were integrated. Pain would be perceived only if the noxious input exceeded the inhibition of pain. However, gate control theory cannot explain the full spectrum of acupuncture effects, and in particular the prolonged pain relief.

Since the 1970s, the secretion of a range of biochemicals or neurotransmitters has been considered among the mechanism of acupuncture analgesia, such as adenosine, opioid peptides, cholecystokinin octapeptide, 5-hydroxytryptamine, noradrenaline, glutamate, GABA, substance P, calcium ions, angiotensin II, somatostatin, arginine vasopressin, and dopamine [69–81].

Different subtypes of opioid receptor were also believed to mediate the frequency-dependent electroacupuncture analgesia [82, 83]. For example, EA at a low frequency of 2 Hz would facilitate the release of enkephalin, but not dynorphin, while a high frequency of 100 Hz would stimulate the dynorphin but not enkephalin release in rats [82], as well as in humans [83]. The primary foundations for acupuncture effects seem to be bioelectromagnetic, while biochemical factors would be secondary. By the way, the opioid peptide secretion was recently proposed as being due to mechanical acoustic shear wave activation and calcium signalling induced by needle rotation [22].

Recently, the inflammatory reflex (via theANS) has been observed as potentially crucial for the antihyperalgesic effect of acupuncture: by regulating the immune system it can elucidate not only the analgesic, but also the anti-inflammatory mechanism of acupuncture [76].

5.9. Acupuncture and Control of Spasticity. Spasticity is a frequent consequence of sABIs, arising from an anarchic reorganization of the pyramidal and parapyramidal fibers, and leading to hypertonia and hyperreflexia of the affected muscular groups and, if untreated, to possible irreversible joint lesions. Current treatment options include intrathecal baclofen and soft splints, botulinum toxin, or cortical activation by thalamic stimulation [84]. There is a low quality evidence for rehabilitation programs, extracorporeal shockwave therapy, transcranial direct current stimulation, transcranial magnetic stimulation, and transcutaneous electrical nerve stimulation targeting spasticity, while a moderate evidence has been shown for electromyomuscular stimulation and acupuncture as an adjunct therapy to conventional routine care (pharmacological and rehabilitation) [85]. In patients with DOCs, acupuncture at GV26, Ex-HN3, LI14, and ST36 was proven to reduce spastic muscle hypertonia by decreasing the excitability of the spinal motor neurons, both ten minutes after needles insertion and ten minutes after their removal [86].

6. Final Considerations and Conclusions

The World Health Organization has recommended acupuncture in 1980 as an effective complementary therapy for several diseases. Among the indications, neurologic disorders have been shown to benefit from acupuncture.

In this review, we analysed scientific studies and clinical reports that explored the acupuncture’s effects in several acquired brain injuries, aiming to

1. limit brain secondary injury, by acting on systemic and local inflammation, oxidative stress, intracellular calcium overload, neuron regeneration, and growth factors release;
2. manage sABI consequences, such as neuroendocrine and autonomic dysfunction, muscle spasticity, and pain.

Research in this field has obtained significant improvement with the technical support of the life sciences, and the studies of acupuncture have in turn accelerated the development of biomedical science. However, intrinsic aspects of this medical approach make it difficult to run a clinical study, and several data derive from animal studies or from small-size and heterogeneous samples of patients. Moreover, the acupuncture selected for treating sABI can differ between research groups.

In addition, patients with disorders of consciousness are per se difficult to study. Given the impossibility of communicating with the patient, the content of consciousness can be only inferred by response behaviour. Diagnostic errors may depend not only on the operator but also on wakefulness fluctuations of the patient, who may be drowsy or agitated, or have epileptic seizures or aphasia. The neurophysiologic evaluations are made difficult by the presence of sweat of the head (which worsens the EEG impedance), muscle hypertonus (that obstructs mobilization), and noise from electromedical equipment, while functional imaging techniques are expensive and not easily available.
### Table 3: Acupuncture and neuroplasticity.

<table>
<thead>
<tr>
<th>MAIN MECHANISMS BY WHICH ACUPUNCTURE MAY INFLUENCE THE PHYSIOLOGIC PLASTIC REACTIONS TO BRAIN INJURY</th>
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<tbody>
<tr>
<td>(i) Inhibition of brain neuronal apoptosis</td>
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<tr>
<td>(ii) Inhibition of aberrant astrocyte activation</td>
</tr>
<tr>
<td>(iii) Upregulation of neurotrophins expression</td>
</tr>
<tr>
<td>(iv) Upregulate expression of GDNF</td>
</tr>
<tr>
<td>(v) Increased functional connectivity</td>
</tr>
<tr>
<td>(vi) Enhanced neuroblast proliferation and differentiation</td>
</tr>
<tr>
<td>(vii) Reduction of blood-brain barrier permeability in intracerebral haemorrhage (caveolin-1/matrix metalloproteinase)</td>
</tr>
<tr>
<td>(viii) Regulation of blood lipid metabolism to counteract cerebral free radical damage</td>
</tr>
<tr>
<td>(ix) Promoting cerebral vascular immunoinflammatory reactions</td>
</tr>
<tr>
<td>(x) Increase of GABA level</td>
</tr>
<tr>
<td>(xi) Reduce contents of excitatory amino acids</td>
</tr>
</tbody>
</table>

**Figure 1: Role of acupuncture in the management of sABIs.**

Further studies are needed to identify the most efficient and customized therapeutical protocol, aiming in particular at eliciting arousal.

The available data suggest that, in patients with sABIs/DOCs, acupuncture may represent an interesting frontier in the years ahead, as it seems to limit the secondary brain injury development, modulate ANS, and ameliorate their quality of life (Table 3, Figure 1). The absence of side effects or drug interactions make it particularly indicated for such fragile subjects.
Conflicts of Interest
The authors declare that they have no conflicts of interest.

References


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