Traumatic Brachial Plexus

Subjects: Clinical Neurology

Contributor: Ilaria Percivale , Massimiliano Leigheb , Stefano Tricca , Andrea Paladini , Giuseppe Guzzardi

Traumatic brachial plexus injuries are rare but serious consequences of major traumas. Pre-ganglionic lesions are considered irreparable, while post-ganglionic injuries can be potentially treated if an early diagnosis is available.

brachial plexus MRI scan MRI diffusion weighted nervous system traumas

peripheral nerves

1. Introduction

The brachial plexus (BP) is the neural network that provides innervation to the upper chest, shoulders, and upper limbs. It is formed by the anterior branches of the last four cervical nerves (C5, C6, C7, and C8) and the first thoracic nerve (T1); the posterior and anterior nerve roots carry, respectively, sensory and motor fibers and exit from the spinal canal through the intervertebral foramen [1].

Before the union of the fibers there is an important structure, the posterior or dorsal root ganglion (DRG), which is considered an important landmark: lesions occurring proximally to DRG are defined pre-ganglionic, while lesions occurring distally to DRG are defined as post-ganglionic.

The second division of the BP is represented by three primary trunks: the superior trunk (formed by the union of C5 and C6 anterior roots), the middle trunk (which is the continuation of C7 anterior root), and the inferior trunk (C8 and T1 roots). The trunks are typically described as running into the interscalene triangle with the subclavian artery [2][3].

Near the lateral border of the first rib, each trunk splits into two branches: anterior and posterior. The six divisions form a triangular cluster that can be identified until the coracoid process occurs, where they form three cords.

The cords—lateral, posterior, and medial—run close to the axillary artery towards the pectoralis minor muscle, where they separate into five terminal branches: the axillary nerve, the median nerve, the musculocutaneous nerve, the radial nerve, and the ulnar nerve ^[1].

Traumatic BP injuries affect 1% of patients involved in major trauma (car accidents, occupational injuries, and falling), causing disability, pain, psychologic morbidity, and reduced quality of life ^{[2][3][4]}.

According to the Seddon, Sunderland, and MacKinnon classifications, traumatic plexopathies can be divided into six degrees based on the number of layers damaged: neuropraxia (first degree), axonotmesis (from second to fourth degree), and neurotmesis (from fifth to sixth degree) ^{[5][6]}.

Neuropraxia is a clinical condition characterized by temporary loss of function without denervation atrophy of the muscle. Axonotmesis is characterized by a Wallerian degeneration followed by nerve regeneration. While the latter can be managed conservatively, neurotmesis needs surgery for axon and myelin sheath disruption ^[Z].

Another important classification of nerve injuries is based on their location: pre-ganglionic lesions are considered irreparable, while post-ganglionic injuries can be potentially surgically treated if an early diagnosis is available. Early surgical nerve repair leads to better functional recovery of the upper limb function ^{[8][9]}.

As a consequence, diagnosis is important to distinguish low-grade lesions not requiring surgical treatment from high-grade lesions and to identify their location ^{[10][11]}. As magnetic resonance imaging (MRI) is a non-invasive, non-radiative imaging modality with multi-planar capability and great soft tissue characterization, it is a basic diagnostic imaging modality ^[12].

Many authors have examined the role of MRI in the diagnosis of traumatic BP injuries.

2. Strategy Search

After searching in the aforementioned internet databases and removing duplicates, 71 articles were retrieved. These studies were then screened for eligibility as presented in the flow-chart (Figure 1). Eight articles underwent a full text screen and four of them were excluded because they were lacking adequate data regarding post-ganglionic BP injuries. Four studies were included in our systematic review, as summarized in Table 1. Of these, three were included in the meta-analysis ^[14][15][16][17], while Caporrino et al. ^[18][18] was excluded from the quantitative synthesis since TP, FP, TN, and FN were not reported in the text. All the included studies had prospective design and considered patients with traumatic BP injuries. All the studies but Caporrino et al. reported the number of patients included ^[15][16][17]. Two of the four studies provided information about the age range of the patients ^[16][17]. In Acharya, Caporrino, and Gad, a 1.5T MRI scanner was employed ^[15][16][18], while in Zhang, a 3T MRI scanner was used ^[17]. All the studies but Caporrino provided a precise description of the employed MRI protocol ^[15][16][17]. All the included studies used surgical findings as standard of reference ^[15][16][17][18].

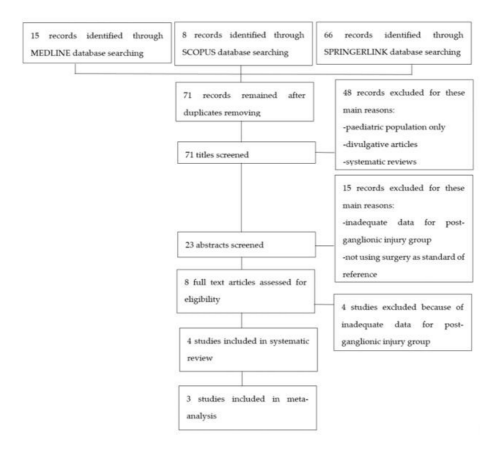


Figure 1. Flowchart of study selection process.

3. Methodological Quality Assessment

Quality assessment of the included studies was conducted with the QUADAS-2 tool (Table 2) ^[11]. All the included studies but Zhang provided adequate information about patients' inclusion and exclusion criteria ^{[15][16][18]}. MRI protocol was extensively described in all the studies but Caporrino ^{[15][16][17]}. It was mentioned in Gad only that surgeons were blinded to the MRI results, and therefore the reference standard was considered unlikely to have introduced biases ^[15]. Only Acharya's article provided clear information about both the time intervals between injuries and MRI and between MRI and surgery; It was then considered at low risk of bias in terms of "flow and timing" ^[16]. Caporrino et al. only reported the time interval between injury and MRI ^[18].

4. Synthesis of Results

Table 1 shows characteristics and main conclusions of the selected studies.

MRI findings considered as significative for post-ganglionic injury of the BP were:

- nerve rupture: characterized by different degrees of nerve thickening caused by edema and inflammation with abnormal hyper intense signal in T2/short-tau inversion recovery (STIR) sequences;
- neuroma formation, characterized by a focal thickening of the injured segment of the nerve $\frac{17}{2}$.

The selected studies did not clearly distinguish data among the different type of lesion.

Table 3 shows sensitivity, which refers to the true positive rate (true positives)/(true positive + false negative), and specificity, which refers to the true negative rate (true negatives)/(true negative + false positive), values with 95% confidence intervals of MRI for traumatic post-ganglionic lesions for each study included in the meta-analysis and the relative forest plots ^{[15][16][17]}.

The paper written by Caporrino et al. was also selected for the systematic review and reported a sensitivity of 60% (95% CI 32.3–83.7) and a specificity of 59.8% (95% CI 48.7–70.1%) ^[18].

The pooled sensitivity, pooled specificity, and 95% confidence interval of the three studies included in the metaanalysis are shown in Table 4. Pooled sensitivity turned out to be of 90% (95% CI 0.78–0.97) and the pooled specificity of 90% (0.86–0.94). The sensitivity value is, however, associated with a l^2 rate >75%, due to the heterogeneous results of the selected literature.

	Study Design	Subject Features	Postganglionic Lesions	Age	MRI Field Intensity	MRI Sequences Employed	MRI Timing	Standard of Reference	Level of Evidence	Main Conclusion
Acharya, 2019 ^[16]	Prospective	35 patients with traumatic brachial plexus injuries	Eight surgically demonstrated postganglionic lesions	Patients under the age of 60	1.5 T	T1-T2-T2 weighted 3D neurography- T2 spin echo- short- tau inversion recovery (STIR)	At least 3 weeks after injury	Surgery	2b	Magnetic resonance imaging (MRI) is a useful tool in the diagnosis of brachial plexus injuries.
Zhang, 2018 ^[<u>17</u>]	Prospective	28 patients with traumatic brachial plexus injuries	23 surgically demonstrated postganglionic lesions, in 12 patients	Mean age: 27.2	ЗT	T1-T2-STIR- balance FFE- diffusion- weighted imaging with background signal	Not reported	Surgery	2b	MRI is a valuable diagnostic tool for brachial plexus lesions, especially if balance-FFE, STIR, and DWIBS

Table 1. Summary of included studies.

						suppression (DWIBS)				sequences are performed.
Caporrino, 2014 ^[18]	Prospective	34 patients with traumatic plexus injuries	Not reported	Mean age: 29.8	1.5 T	Not reported	2–3 months after injury	Surgery	2b	MRI showed poor diagnostic performance in identifying brachial plexus lesions compared to physical examination. Notwithstanding, it is reasonable to think that the combination of physical examination and MRI could provide the best diagnostic accuracy.
Gad, 2020	Prospective	22 patients with traumatic brachial plexus injuries	18 surgically demonstrated postganglionic lesions	Mean age: 26.3	1.5 T	T1, STIR, T2, T2-STIR, and DWIBS	Not reported	Surgery	2b	"MRI is the imaging modality of choice in the examination of traumatic and obstetric brachial plexus injuries; it is safe and non-invasive, having the multiplanar capability

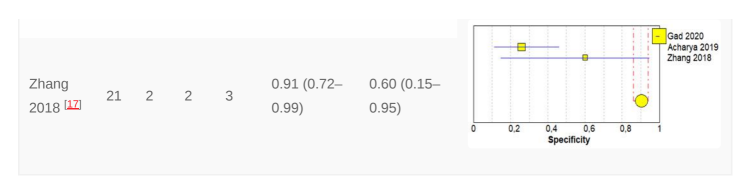
and better soft tissue characterization".

Table 2. Quality assessment of diagnostic accuracy studies (QUADAS)-2, quality assessment of the included studies.

	Patient Selection	Index Test	Reference Standard	Flow and Timing
Acharya, 2019 ^{[<u>16]</u>}	+	+	+	+
Zhang, 2018 [<u>17</u>]	?	+	?	?
Caporrino, 2014 ^[<u>18</u>]	+	?	+	?
Gad, 2020 ^[15]	+	+	+	?

Table 3. Forest plot showing sensitivity and specificity for each included study.

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Forrest Plots
Gad 2020 ^[<u>15</u>]	16	0	2	198	0.89 (0.65– 0.99)	1.00 (0.98– 1.00)	Gad 2020 Acharya 2019 Zhang 2018 0 0.2 0.4 0.6 0.8 1 Sensitivity
Acharya 2019 ^{[<u>16]</u>}	7	20	1	7	0.88 (0.47– 1.00)	0.26 (0.11– 0.46)	



Abbreviations: TP, true positive; FP, false positive; FN, false negative; TN, true negative.

		Pooled Sensitivity		Pooled Specificity		Pooled LR+		Pooled LR-		Pooled DOR			
TP	FP	FN	TN	Value (95% CI)	l ²	Value (95% CI)	²	Value (95% CI)	²	Value (95% CI)	²	Value (95%Cl)	²
44	22	5	208	0.90 (0.78– 0.97)	0.0%	0.90 (0.86– 0.94)	98.1%	7.70 (0.28– 214.76)	96.5%	0.17 (0.07– 0.39)	0.0%	40.71 (0.99– 1666.3)	84.6%

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Abbreviations: TP, true positive; FP, false positive; FN, false negative; TN, true negative; CI, confidence interval; LR+, positive likelihood ratio; LR-, negative likelihood ratio; DOR, diagnostic odd ratio.

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