

CCL-2 as a potential marker for remission after traumatic spinal cord injury in Chinese patients.

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Abstract

Objectives: To state the correlation between CCL-2, CCL-4 and CXCL-5 serum degrees and remission following traumatic spinal cord injury (TSCI) in a human protocol in comparison with animal research works.

Methods: We performed the examination of the serum levels of CCL-2, CCL-4 and CXCL-5 across an eight week duration; especially, at the time of admittance and 4, 9 and 12 h, 1 and 3 days and 1, 2, 4, as well as 8 weeks subsequent to trauma. In accordance with the design of our research, we performed a matching 12 Chinese patients with TSCI as well as neurological remission with 12 Chinese patients having a preliminary ASIA A grade together with no neurological remission. Overall 12 patients having vertebral fracture without neurological shortages acted as control. Our analysis was carried out with the use of a Luminex Cytokine Panel.

Results: The observation received through our research exhibited dissimilarities in the serum expression layouts of CCL-2 in relation to the neurological remission (CCL-2 at admission $P=0.001$). Serum degrees of CCL-2 as well as CCL-4 were considerably varied in patients having or not having neurological remission.

Conclusion: As indicated by our findings, peripheral serum analysis is considered as an appropriate idea for the prediction of the patient's capacity for neurological remission subsequent to TSCI. That is why the current research work provides the introduction of a potential technique for prospective inspection of the ideas together with tracking methodologies for the prevalent therapies. As indicated by the findings, prospective examinations with an extended specimen size are required for the purpose of developing inspection, prognostic as well as scoring systems.

Keywords: Spinal cord injury, CCL-2, Inflammatory factor.

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Introduction

Spinal cord injury (SCI) continues to be one of the most demanding injuries in the contemporary medicine. SCIs together with consequent paraplegia sternly bothers patient's standard of life, in addition to giving rise to fiscal issues for not only the patient but also the associated families [1-3]. In addition to surgical treatment as well as rehabilitation, as of now, no direct treatment for traumatic SCI, for instance medication, is existent, accordingly constraining the success of accessible cures and letting an elevated threat of a discontented result [4,5]. In addition, proof in respect of the most therapeutic strategies is also very rare [6]. Till today, no authentic marker is available that is capable of indicating a promising capacity for the remission in Chinese patients. That is the reason additional information on the biochemical mechanisms is required following the SCI. The pathophysiological mechanism following the SCI is considered as intricate, together with comprising a first and second stage, which are associated with the mechanical trauma and,

thereafter, an array of pathophysiological mechanisms. All through the sub-acute phase, the inflammatory feedback counts on a preliminary wave of infiltrating neutrophils, which precede a shift of monocytes as well as macrophages into the damaged area with the use of several processes.

In this research work, we aimed at determining a prognostic framework in order to evaluate the remission capacity following SCI on the bases of promising biomarkers CXCLs as well as CCLs in peripheral blood serum [7]. Because of their significance in (neuro-) inflammatory mechanisms, especially subsequent to the traumatic damages, we made a decision to perform the investigation of the serum degrees of CCL-2, CCL-4 as well as CXCL-5 [8-10]. The findings of this research aim at providing an advanced possible diagnostic approach for helping predict the potential remission following SCI, in addition to assessing the clinical therapies in respect of their achievements on remission levels [11-12]. That is why this research threw focus on the following research questions: Is it probable to make prediction of the rehabilitation subsequent to

SCI on the bases of serum cytokine expression levels? Is it likely to make use of analysis of the serum chemokine expression levels as a latent diagnostic instrument to monitor the neurological remission in prospective treatments subsequent to SCI? The feedbacks to these questions are expected to assist us in getting closer to the monitoring enhancement, together with establishing a purposeful score in the prediction of the remission following SCI.

Materials and Methods

From the year 2012 to 2016, 120 patients (93 men and 27 women) related to TSCI together with having received treatments in our hospital were added in our register in a prospective manner. Acquisition of blood specimen occurred in every scenarios featuring a time precision of $\pm 10\%$: 4 vials of serum (each 7.5 ml) were attained at 4, 9 and 12 hours, 1 and 3 days as well as 1, 2, 4, and 8 weeks subsequent to SCI. Following twenty minutes coagulation, centrifugation of the blood specimens was performed at 3000 rpm, followed by aliquotion and storages at minus 80 °C until analysis.

In accordance with the corresponding status of the database, analyses of varied gatherings of serum specimens have been performed for addressing corresponding scientific problems [13-16]. Evaluation of the AIS grades occurred at the entry, in addition to, subsequent to twelve weeks as per the International Standards for Neurological Classification of SCI (ISNCSCI). The definition of the criterion regarding neurological remission was providing as favorable transformation of AIS grades within 12 weeks following trauma. All ISNCSCI inspections were carried out by the major physical therapist. Preliminary ISNCSCI inspections were carried out within the period of first 72 hours following the entry. Admissions of the patients employed for the registration were done within 2 h following the activation of the emergency response system. Realization of surgical therapy, thereafter, was performed within 3.5 ± 1.6 h following trauma. The GLP (Good Laboratory Practice) deliveries established the standard for all of the inspection of serum specimens. The use of Luminex Performance Human High-Sensitivity Cytokine Panels was made for the purpose of the quantitative determination of CCL-2, CCL-4 and CXCL-5 in the patients' serum specimens. Determination was carried out in accordance with the guidelines of the manufacturer (Catalog Number FCST03-04, Kit Lot Number 1420835). R&D Systems (Minneapolis, MN, USA) delivered the kits. The corresponding analytes' attributes have been provided hereunder in sensitivity (pg ml⁻¹) and standard curve range (pg ml⁻¹): CCL-2 (9.9; 98–7940), CCL-4 (5.8; 261–21 150) and CXCL-5 (8.2; 50–12 140). Performance of all of the quantitative analyses was done in Union Hospital having affiliation with the Tongji Medical College by a lab personnel who wasn't aware of patients as well as medicinal information.

Approval of the recent research has been received from the ethics committee of the Huazhong University of Science and

Technology. Willful signing of the consent forms, together with their dating, was done by all of the research participants. All of the patients were able to make volunteer choices regarding quitting the research at any moment due to any issue. The patients that matched with the ruling out criteria (nontraumatic spinal cord injury, traumatic brain injury (TBI), severe abdominal trauma, traumatic amputation of extremities, coma and additional major trauma apart from the SCI) were not included without any omission. There was no patient that had been administered methylprednisolone sodium succinate (MPSS) all through the research contribution. Matching

For the purpose of eliminating the medicinal covariates on the corresponding cytokine level, cohorts of corresponding pairs were formed for the purpose of investigating the dissimilarities within their cytokine layouts. That is why we corresponded 12 patients with SCI and neurological remission (Group 1=G1) with 12 patients having a preliminary American Spinal Injury Association (ASIA) A grade together with no neurological remission (Group 0=G0). In addition, 12 patients having a remote vertebral fracture with no neurological shortages acted as a control group (Group C=C).

Matching criteria involved the factors hereunder: age, sex, etiology and AO (“Arbeitsgemeinschaft für Osteosynthesefragen”) fracture classification. Provided the situation that one plus matches were able to be allocated, and then the patient having the most comparable medical status was given preference and vice versa. The consequent patients' demographics have been presented in Table 1, suggesting the AO classification of every patient in group C. The matching mechanism was carried out with a retrospective approach on the bases of the medical data. Patient registration, specimen gathering together with analysis of CCL-2, CCL-4 and CXCL-5 were carried out prospectively.

Statistical analysis

Performance of explorative correlation analyses was done among all variables. Evaluations of the nonparametric test approaches were done for the purpose of investigating the location transfers between (Mann–Whitney U-test) and within groups at varied moments (Wilcoxon Signed Rank Test). Analysis of 2 plus sovereign specimens were carried out with the use of Kruskal–Wallis test. Evaluations of the categorical variables were performed with the use of the χ^2 test.

Results

The design of the current research follows a prospective as well as explorative medical investigator research. Patients involved in this research matched out of 120 situations in our database. No randomization was observed. Criteria to perform the match-pair analysis possessed the patient's sex, age, etiology as well as AO categorization (Table 1). Documentation of the patients' demographics was performed;

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analyses as well as comparisons of the groups were carried out as formerly published [15-18].

Table 1: a) Demographic and clinical characteristics of patients with TSCI and b) referring to the control group C.

(a) Patients	Remission	No remission	t/ χ^2	Comparison G1 G0
	(G1)	(G0)		(P-value)
N	12	12		
Age (years)	43 ± 12	45 ± 23	0.267	0.792
Time accident/alarm to admission (min)	73 ± 22	72 ± 32	0.089	0.93
Gender				
Female	4	3		1.000
Male	8	9		
Etiology				
Fall	5	4		1.000
Traffic	7	8		
AO classification				
A	7	8		1.000
B	3	2		
C	2	2		
NLI				
C	3	6		0.726
Th	4	3		
L	3	1		
None	2	2		
AIS at admission				
A	9	10		
B	2	2		
C	1	0		
D	0	0		
E	0	0		
AIS after 12 weeks				
A	0	10		
B	1	2		
C	5	0		
D	6	0		
E	0	0		
(b) Patients	Control (C)			
N	12			
Age (years)	43 ± 12			
Gender				

Female	6
Male	6
Etiology	
Fall	8
Traffic	4
AO classification	
A	9
B	3
C	0

Age has been stated as mean years \pm s.d. Definition of neurological remission was made as enhancement in AIS. Performing the comparison of G0 and G1, analysis of P-values was conducted with the application of either the χ^2 test or Mann-Whitney U-test, whereby the exhibit dissimilarities between G0 and G1.

Patients' demographics

In accordance with our developed matched-pair analysis, 36 patients were involved within this research work (13 women and 23 men) out of our current database that includes 120 scenarios, which were given admission from the year 2012 until the year 2016. The average age amounted to be 43 ± 16 years. The study group (Group S) comprised twenty four patients that had neurological impact of TSCI. The control group (Group C) comprised twelve patients that had received a traumatic damage with no neurological injury. In respect of group S, there were twelve that had (Group 1 remission) and twelve that didn't have AIS damage (Group 0 no remission). The ones having no AIS injury were selected for having a preliminary AIS grade A. In C, no incisions were observed on the spinal cord. All of the medicinal attributes have been presented in Table 1.

Statistically, no dissimilarities were observed between research groups in respect of age, duration from accident/alarm to entry, period of the 1st operation, gender, etiology as well as AO categorization (termed as G1 and G0 in the following chapter). All of twenty four patient that had traumatic SCI (G0 and G1) underwent surgery (12 ventral 50.0%; 12 dorsal 50.0%); 15 were given spondylodesis treatment (75.0%). All of thirty six were added in the combined suffered vertebral fractures. The AIS grades at both the entry and discharge together with the NLI and the kind of paralysis were considerably varied in both G1 and G0 ($P < 0.05$).

CCL-2

Within the initial twelve hours, average CCL-2 values amounted to be higher in G0 as compared with that in G1 and C. Levels of serum in G0 dropped from 699.36 ± 104.54 pg ml⁻¹ to 429.45 ± 52.50 pg ml⁻¹ 4 h subsequent to the trauma. In respect of G1, average CCL-2 values exhibited a minimum 288.04 ± 67.50 pg ml⁻¹ at 12 hours, together with top serum levels 456 ± 67.50 pg ml⁻¹ 24 hours following the trauma. Levels of serum in G1 stood lower as compared with that in G0 all through the duration, other than for the duration of top levels in G1 24 hours following the trauma. In respect of group

C, average CCL-2 levels amounted to be lower as compared with those of G0 and G1 all through the entire time period; serum levels displayed a reducing tendency from preliminary 349.20 ± 95.80 pg ml⁻¹ to 247.13 pg ml⁻¹, 1 week subsequent to the trauma. Dissimilarities between G0 and G1 at admission ($P = 0.001$) as well as 2 months ($P = 0.039$) subsequent to trauma were quite considerable as their analysis was performed with the application of the Mann-Whitney U-test. In addition to that, findings of our research exhibited a considerable preliminary height of CCL-2 in G1 in comparison with that in G0 ($P = 0.026$).

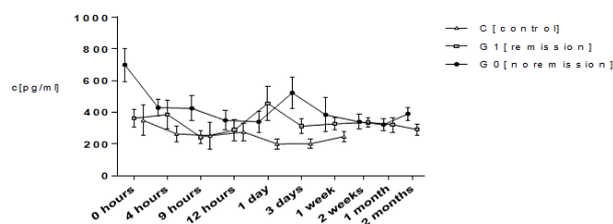


Figure 1: Serum expression patterns of G0, G1 and C expressed as means \pm s.e. of the mean and Serum level comparison of G0 and G1 of CCL-2 concentration. Differences were assessed with the Mann-Whitney U-test between both groups at each particular time point.

CCL-4

Average CCL-4 values of patients took start ranging between 108.79 and 175.95 pg ml⁻¹. That of G0 and C continued exhibiting a steady level. At a point of time of nine hours subsequent to trauma in G1, CCL-4 values showed a rise from 78.03 ± 17.59 pg ml⁻¹ to the topmost of 214.20 ± 96.34 pg ml⁻¹ at twenty four hours that dropped once again to 96.18 ± 19.13 pg ml⁻¹ 1 week subsequent to trauma. Average levels of serum of G0 as well as G1 emerged falling between the ranges of 50.49 and 97.92 pg ml⁻¹ subsequent to two months. Moreover, the average levels in C continued to be on a steady level of 38.72 ± 13.25 pg ml⁻¹ 1 month subsequent to trauma. In respect of group C, average CCL-4 serum levels amounted to be lower in comparison with those of G0 and G1 always. Analyzing them using the Mann-Whitney U-test brought forth

considerable dissimilarities nine hours ($P=0.037$) and 2 months ($P=0.043$) subsequent to trauma between G0 and G1. Average levels in G1 displayed a rise to the topmost at twenty four

hours, suggesting higher levels for providing an advantageous impact on TSCI remodeling mechanisms.

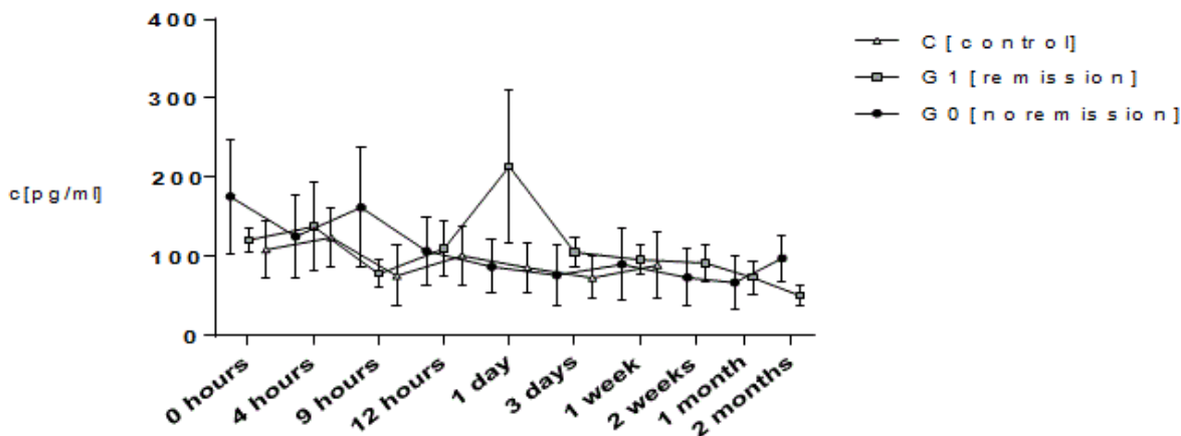


Figure 2: Serum expression patterns of G0, G1 and C expressed as means±s.e. of the mean and Serum level comparison of G0 and G1 of CCL-4 concentration. Differences were assessed with the Mann–Whitney U-test between both groups at each particular time point.

CXCL-5

The expression layout of average CXCL-5 values of all of the subgroups G0, G1 and C exhibited an identical layout. Taking a stat between the ranges of 713.30 and 1038.20 pg ml⁻¹, all of them touched their topmost two weeks subsequent to trauma in the ranges between 1637.15 and 1935.45 pg ml⁻¹. Ultimate average values in both G0 and G1 following two months (G1: 1252.90 ± 201.025 pg ml⁻¹; G0: 1112.65 ± 392.70 pg ml⁻¹) and in C following two weeks (C: 1637.15 ± 319.73 pg ml⁻¹) amounted to be higher in comparison with that in the start (G1: 1038.20 ± 210.38 pg ml⁻¹; G0: 713.3 ± 280.50 pg ml⁻¹; C: 925.65 ± 219.73 pg ml⁻¹). Analyzing them using the Mann–Whitney U-test exhibited immaterial dissimilarities between groups G0 as well as G1 always.

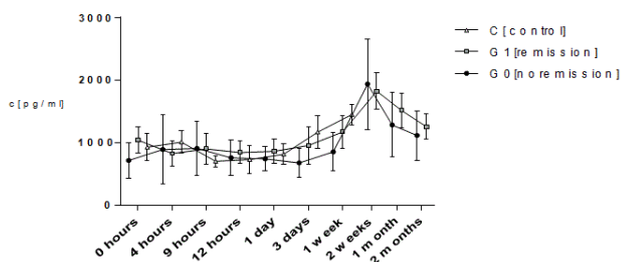


Figure 3: Serum expression patterns of G0, G1 and C expressed as means±s.e. of the mean and Serum level comparison of G0 and G1 of CXCL-5 concentration. Differences were assessed with the Mann–Whitney U-test between both groups at each particular time point.

Discussion

In this research, we aimed at determining a prognostic framework in order to evaluate the remission capacity subsequent to after SCI on the bases of promising biomarkers CXCLs as well as CCLs in peripheral blood serum. The

observations of our research bring forth material information regarding the preliminary evaluation of TSCI patients. As suggested by our data, dissimilarities of CCL-2 between G1 and G0 at entry (362.13 ± 55.55 vs. 699.36 ± 104.54 ; $P=0.001$) as well as CCL-4 at nine hours (78.03 ± 17.59 vs. 162.18 ± 75.32 ; $P=0.037$) imply those patients highly prone to a neurological injury within the first four hours following trauma. Moreover, the observations of our study provide the introduction of a predictive value for the distinction between patients having no probability of neurological remission and those having the elevated chances for remission.

Strength and Limitations

On the bases of future investigative research design with matched pairs, we performed the investigation of the expression layouts of CCL-2, CCL-4 as well as CXCL-5 in human peripheral serum all through the severe, subacute and mid-level stage of traumatic SCI in accordance with a trusted protocol. Our extensive database ($n=120$) made us able to carry out a precise match-pair analysis. Despite the fact that the findings from animal frameworks are constrained by their ability to be transferred [11], yet the limitless specimen size continues to be beneficial. The current research work added a tiny group of 36 patients of matched pairs, and, that is why findings are not likely to be the representative of domestic mechanisms obviously. We made a decision for the inclusion of those patients that had varied degrees of spinal cord injury together with varied categorizations in the ASIA score in the remission cohort. In respect of the no-remission group, merely ASIA A patients was involved, doing away with the partiality of probable confusing information. Additional research works with more extended specimen size are required for addressing an inter-group (G1 vs. G0) comparison with each AISA grade, owing to the fact that the prognosis is mostly different by a

sizeable degree. Owing to the huge standard errors of the average as well as the petite numbers of patients, the sizeable dissimilarities as stated are mandatorily required to be taken into consideration in more of a careful manner. For the purpose of further investigating the function of CCLs as well as CXCLs in the pathophysiology following TSCI, they are required to be associated with the developed inflammation markers, for instance Interferon- γ , Tumor necrosis factor- α , Interleukin-1 β , IL-6, IL-8, IL-10 as well as vascular endothelial growth factor. In addition, we were capable of presenting discrete expression layouts in respect of both CCL-2 and CCL-4, suggesting their implication in the second stage of injury regarding G0 and G1. The two also bring forth additional information regarding the demarcation between S and C.

CCL-2: CCL-2 (termed as monocyte chemoattractant protein-1 MCP-1 as well) is quite popular for being heightened in the CNS and cerebrospinal fluid of individuals having neuroinflammatory signs typified by blood-brain barrier (BBB) disruption as well as leukocyte penetration of the CNS [19,20]. That is why we develop a presumption that CCL-2 is to be considered as well in continued neuroinflammatory mechanisms following the traumatic SCI. In the year 2012, Roberts et al. [21] suggested that CCL-2 triggers shuttling of β -catenin between PECAM-1 (platelet endothelial cell adhesion molecule-1) as well as the adherens junction (AJ) in human brain microvascular endothelial cells in a way that the AJ and monolayer integrity receives transient disruption.

For the purpose of gaining more peculiar together with precise models, there is a need to perform additional research works with an extensive collective. Owing to the constrained specimen size, the current framework is likely to receive the impact from overfit. That is why we make strong recommendation to perform cross-validation on framework in prospective researches with an extensive collective for avoiding the partiality.

CCL-4: Being the major mediators of bond to the endothelial top, chemokines perform a key function in the employment of T-cell subsets (formerly confined to the CNS in vivo) across the BBB all through the CNS inflammation. Observation made by Quandt et al. [22] in the year of 2010 threw emphasis on the function and significance CNS-derived CC-chemokines for the regulation of the traffic of currently activated T-cell subsets across cytokine-activated cerebral endothelium in inflammatory ailments. They stated CCL-4 together with CCL-5 for enhancing the bond of particular CD4+ T-Cell subsets to human brain endothelial cells. There is need to perform additional research for the purpose of clarifying reasons regarding the substantial CI in G1 at twenty four hours. Latent details are likely to be attained with the help of categorization and phenotypisation of immune cells in the incision site, followed by complementation by a functional inspection. Correlations of CSF, Serum and parenchymal cytokine expression are expected to deliver further information regarding the comprehension of biochemical mechanism, together with the establishment of forecasting and inspecting systems.

CXCL-5: Despite the reports suggesting the CXCL-5 possessing neuroprotective impact subsequent to TSCI [23], yet the findings of our research throw light on no additional information in respect of neither recognition of a continued neurotrauma nor distinction between G1 and G0. CXCL-5 somewhat suggests a systemic implication in overall recovery mechanisms subsequent to trauma. Additional research, that is why, is required to throw emphasis on CXCL locally.

Conclusion

Data for systemic degrees of CCL-2 and CCL-4 extracted from peripheral serum analysis bring forth critical information regarding the biochemical mechanisms continued subsequent to TSCI. Despite the fact that critical consideration should be given to the current findings because of petite patient collective, yet they are likely to develop the foundation of prospective study dealing with the investigate as well as extrapolative scoring methodologies in TSCI with the use of research with extended specimen size.

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