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1	Study of Neuron-Specific Enolase as Potential Biomarker for
2	Assessing the Severity and Outcome in Patients with
3	Cerebrovascular Accidents
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8 Abstract

- 9 Background: Stroke is the third cause of death and foremost cause of disability worldwide.
- ¹⁰ Cerebrovascular accident or stroke is an emergency condition which require immediate
- ¹¹ procedure by a neurologist. Determination of extent of brain damage at the onset of the
- ¹² seizure is an appropriate action to determine therapy and prognosis. Increased serum neuron
- ¹³ specific enolase can be expected to differentiating stroke types at the onset of the seizure. Aim:
- ¹⁴ The goal of present study was to measure diagnostic value of serum neuron specific enolase in
- ¹⁵ various types of cerebrovascular accident as well as to evaluate the clinical performance of
- ¹⁶ neuron specific enolase in early diagnosis of cerebrovascular accident.
- 17

Index terms— cerebrovascular accident, ischemic stroke, hemorrhagic stroke, neuron specific enolase, brain
 damage.

²⁰ 1 Conclusion:

21 Present study shows that, assessment of serum neuron specific enolase level may be a useful Marker for severity 22 in cerebrovascular accident like ischemic stroke and hemorrhagic stroke and it may be well correlated with 23 neurological disability and short term functional outcomes.

24 2 Introduction

troke or cerebrovascular accident (CVA) is the third leading cause of death after cardiovascular disease and cancer
(1). In fact it is a leading cause of morbidity and mortality in major industrial countries. Approximately 20
million people each year are suffer from stroke (2).

India will face enormous socio-economic burden to meet the cosis of rehabilitation of stroke victims. Because the population, it is now surviving through peak years (age 55-65 years) of occurrence of stroke (3).

The two major mechanisms causing brain damage in stroke are ischemia and hemorrhage. The effects of 30 ischemia are fairly rapid because the brain does not store glucose, the chief energy substrate and is incapable 31 of anaerobic metabolism. Intracerebral hemorrhage originates from deep penetrating vessels and causes injury 32 33 to brain tissue by disrupting connecting pathways and causing localized pressure injury. 4 The diagnostic and 34 management of CVA is limited by lack of rapid diagnostic assay for use in emergency setting. In recent years, 35 neurobiochemical markers of brain damage have gained particular attention in the identification of stroke patients with an adverse neurological outcome. The serum Neuron -Specific Enolase (NSE) level is one of these markers 36 which can provide early information about neuronal damage. 5 37

Neuron -specific Enolase (NSE; EC 4.2.1.11) is an acidic soluble protein which functions as glycolytic isoenzyme. It is a 78 kD gamma homodimer and represents the dominant enolase isoenzyme which found in the cytoplasm of neurons and cells with neuroendocrine differentiation. 6 ischemia and traumatic head injury provides a reliable laboratory indicator of the degree of brain cell damage and may allow for early prediction of 42 outcome. 7 An increased NSE concentration in blood has been reported in patients with small cell lung cancer,
 43 neuroblastoma and neurological disorders. 7, 8

Thus, in view of above information and several risk of complication, it is worthwhile to study the various biomarkers in CVA. Very few studies have been reported from serum NSE testing and its application in Indian

46 context. The initial aim of our study was to measure the serum NSE levels in various types of CVA within in 47 72 hours of admission. Furthermore, the remarkable intention of present research was to evaluate the clinical

48 performance of NSE in early diagnosis of CVA and monitoring tool for early prediction of ischemic stroke.

49 **3 II.**

50 4 Material and Method

The present diagnostic case-control study was conducted at department of Biochemistry in PDVVPF's Medical College Ahmednagar with all participants providing informed consent and utmost care was taken during experimental procedure according to the declaration of Helsinki 1975. hours of admission to exclude patients with stroke mimic. Approximately 5 ml blood was collected by venipuncture from anticubital vein of the forearm of each subject in plain vaccutainer (yucca diagnostic) under aseptic conditions within 72 hrs after admission and centrifuged for serum collection. Serum was stored at 20 0 C until assay was run to evaluate. All samples were thawed and analyzed in a single series.

58 **5** III.

Method 1) Determination of serum NSE: Serum NSE was measured with commercially available quantitative enzyme linked immune sorbent assay (ELISA) kit which based on biotin double antibody sandwich technology. Add serum containing NSE to well that is pre-coated with NSE monoclonal antibody and then incubate. After incubation, add NSE antibodies labeled with biotin to unite with streptavidin-HRP, which forms the immune complex. Remove unbound enzymes after incubation and washing, then add chromomegnic reagent A and B.

64 colour change blue to yellow with effect of acid which positively correlated with concentration of human NSE. 9, 65 10

Statistical analysis: The statistical analysis was carried out by using the SYSTAT software package for window 66 version 12. The students "t" test was applied for the statistical analysis and the results were expressed in 67 mean \pm Standard Deviation (mean \pm SD).p values p<0.05 for NSE were considered to be statistically significant. 68 The Receiver Operating Characteristic (ROC) curve analysis and the area under the curve were performed for 69 determination of diagnostic performance of serum NSE in the all patients included in the study. The optimum 70 cutoff values for determination of serum NSE were selected from ROC analysis. This optimum cutoff was used 71 to dichotomously classify positive or negative serum NSE level, and used for calculating of diagnostic sensitivity 72 and specificity. 73

74 IV.

75 6 Results

Baseline demographic and clinical characterization of the patients and healthy controls groups are given in table-1
 there were no significant differences between the groups in age, gender.

78 7 Discussion

79 Stroke causes a vast amount of death and disability throughout the world. It is important to have sufficiently 80 sensitive marker for brain damage that can be determined in blood instead of cerebrospinal fluid because blood samples can be taken more frequently and more independent of raised intracranial pressure than cerebrospinal 81 samples. 11 Evaluation of enzyme level in Cerebrospinal fluid or serum has evoked keen interest as a simple, 82 economical, reliable and easily available method for the evaluation of severity, course and prognosis and to some 83 extent in the differential diagnosis of various types of CVA.Ischemia causes a cascade of event that eventually 84 leads to neuronal damage and cell death. 12 NSE is the predominant enolase found in neural tissue, and the 85 structural characteristics of this enolase allow for greater stability in high chloride concentrations compared with 86 enolase in other organ system. 7 87

In the current study, Serum neuron specific enolase levels were increased significantly (p < 0.05) in the ischemic 88 stroke group and hemorrhagic stroke as compared to controls group. Our results are strongly supported to 89 90 previous reports. 5,11,13, Increased NSE level in CVA may be due to cute CNS such as cerebral infarction, 91 hypoxia, trauma and seizure the blood -brain barrier is altered and astroglial disintegration results in leakage of 92 NSE into the serum and cerebrospinal fluid. 13 Schaarschmidt H et. al. research where they have studied the 93 NSE in relation to the severity CVA. They verified that, plasma NSE level is seen as a relevant parameter for assessing the prognosis of cerebral ischemia. Additionally it may prove to be a useful tool for monitoring space 94 occupying brain infarctions and intracerebral hemorrhage and therefore may contribute to improved therapeutic 95 management of severe cerebrovascular diseases. 11 96

Numbers of researchers have focused on the study of NSE in various types of CVA. Aparna Pandey et al have
 showed that initial serum NSE level may be a useful marker for severity in acute ischemic stroke and it may be

well correlated with neurological disability and short term functional outcomes. They also suggested that, serum 99 NSE may be used as an indicator of outcome in cerebral infarction patients. 14 Natheer H. Ravi and Karim 100 M. Aantiyah have demonstrated that, salivary NSE alone or in combination with serum can be used as valuable 101 102 diagnostic for measurement of neuronal damage in patients with stroke and stroke related diseases. According to them, in ischemic stroke, the integrity of blood-brain barrier is disrupted to various degrees in these patients 103 and leakage of this enzyme outside the CNS can be seen in salivary secretion. 13 Takaaki kirino et al have 104 documented that, NSE as a reliable enzymatic indicator of axon injury, regeneration and in particular of target 105 innervations and reinnervation. 6 106

In cross-sectional comparative study, Diwi L Lukas et al have examined and compared serum NSE level in 107 ischemic and hemorrhagic stroke patients according to lesion volume and also analyzed correlation between serum 108 NSE level and lesion volume in CT scan as gold standard. According to them, serum NSE level in acute stroke 109 patients (24-48 hours) after onset can be used to estimate the extent of brain damage (lesion volume) but it 110 cannot be used to differentiate the type of stroke. 15 In present study, furthermore diagnostic performance of 111 Serum NSE for diagnosis of ischemic stroke and hemorrhagic stroke were analyzed. The optimum diagnostic cut 112 off point maximizing the sensitivity and specificity was determined to be 40 ng/ml with a sensitivity of 87.10 %113 and specificity 95% the area under ROC curve for NSE was 0.84. Our results are completely conformity with Hill 114 115 et al study. They found that in a single examination, NSE had sensitivity of 89%. They also suggested that, in 116 the future the examination of neurobiochemical marker panel like serum NSE, Myelin basic protein, Protein S-100 117 B etc can be used not only to differentiate the type of stroke but also to differentiate the subtype of acute stroke. 16 In addition, Natheer H Rawi and Karim M Atiyah have accomplished the diagnostic performance of NSE in 118 patients with ischemic stroke and stroke prone patients. According to their result, the area under the ROC curve 119 for serum NSE was significantly higher (0.960) compared to salivary NSE (0.825). The optimum cut-off value 120 for serum NSE showing the highest diagnostic accurancy (90%) was ? 13.1?g/L. This cut-off threshold showed 121 optimum specificity (100%) and reasonable sensitivity (85%). 122

123 **8 VI.**

124 9 Conclusion

 125 $\,$ Our finding indicate that, serum NSE as brain biomarker in CVA had elevated compared to controls which

provides insight into the pathophysiologic mechanism of brain injury. Serum NSE in CVA patients after onset (<72hours) may be used to estimate the extent of neuronal damage and can be reliable parameter for weigh up CVA.



Figure 1:



Figure 2: Figure 2 Figure 1 :

128

Results: serum neuron specific enolase (p<0.05) concentrations was significantly higher in cerebrovascular accident than healthy controls. Sensitivity, specificity, positive predictive value and negative predictive value of neuron specific enolase for detection of cerebrovascular accident were 87.10%, 95.00%, 92.74% and 87.69%. The area under the receiver operating characteristic curve of neuron specific enolase in cerebrovascular accident was 0.84.

[Note: Aim: The goal of present study was to measure diagnostic value of serum neuron specific enolase in various types of cerebrovascular accident as well as to evaluate the clinical performance of neuron specific enolase in early diagnosis of cerebrovascular accident. Methods: A diagnostic case control study was conducted on 60 patients were admitted within 72 hours of onset of stroke in the department of neurology and department of medicine of PDVVPF's Medical College and hospital Ahmednagar and 60 healthy age and sex matched volunteers formed the control group. Serum neuron specific enolase level was estimated by commercially available quantitative enzyme linked immune sorbent assay (ELISA) kit which based on biotin double antibody sandwich technology. Statistical analysis used: The student t test was used to compared patients and control. Receiver operating characteristic curve for neuron specific enolase was established to determined cut-off point. The sensitivity and specificity of neuron specific enolase for detection of cerebrovascular accident were analyzed.]

Figure 3:

			CVA
	Variables $ontrols schemic stroke (n=32)$		
	(n=60)		stroke
			(n=
			(28)
Age in years	$40.1\pm$	41.3 ± 14.06	$43.1\pm$
	12.34		13.01
Gender (Men/Women)	28/30	19/13	11/27
Systolic (mmHg)	blo quitesis10 ;e	$119.15 \pm \ 30.11$	120.07
	±		\pm
	15.03		35.16
Diastolic blood (mmHg)	Pre 551162	81.42 ± 14.28	79.53
	±		±
	5.04		16.02
Cigarette Smokers (n)		07	15
Tobacco Chewing(n)		26	19
Atrial fibrillation (n)	07	06	04
DM (n)	9	12	8
Serum NSE (ng/ml)	$14.55\pm$	$43.63 \pm 13.41^*$	$45.63\pm$
	12.41		15.89^{*}
Values were expressed in mean with Sta	andard Deviation	$(\text{mean}\pm\text{SD}),$	

*Statistically highly significant,(p<0.001)

n =numbers

As shown in table-1, Serum neuron specific enolase levels were increased significantly (p<0.05) in the ischemic stroke Group (43.63 \pm 13.41) and hemorrhagic stroke (45.63 \pm 15.89) as compared to controls group (14.55 \pm 12.64). The performance of Serum NSE for diagnosis of

ischemic stroke and hemorrhagic stroke is presented in

table-2. Sensitivity, specificity, positive predictive v and negative predictive value were 84.38%, 95.00%, 90.00% and 91.94% respectively in Ischemic stroke. Similarly, in hemorrhagic stroke, Sensitivity, specifi positive predictive value and negative predictive vawere 89.29%, 95.00%, 89.29% and 95.00%.

Figure 4: Table 1 :

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۷		2	

Types of CVA	Sensitivity (%)	Specificity (%)	Positive Predictive Value (%)	Negative Predic- tive Value (%)
Ischemic Stroke	84.38	95.00	90.00	91.94
Hemorrhagic stroke	89.29	95.00	89.29	95.00

Figure 5: Table 2 :

9 CONCLUSION

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