Mapping Brain Injury with Symmetrical-channels’ EEG Signal Analysis – A Pilot Study

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A technique for detecting brain injury at the bedside has great clinical value, but conventional imaging techniques (such as computed tomography [CT] and magnetic resonance imaging) are impractical. In this study, a novel method—the symmetrical channel electroencephalogram (EEG) signal analysis—was developed for this purpose. The study population consisted of 45 traumatic brain injury patients and 10 healthy controls. EEG signals in resting and stimulus states were acquired, and approximate entropy (ApEn) and slow-wave coefficient were extracted to calculate the ratio values of ApEn and SWC for injured and uninjured areas. Statistical analyses showed that the ratio values for both ApEn and SWC between injured and uninjured brain areas differed significantly (P<0.05) for both resting and name call stimulus states. A set of criteria (range of ratio values) to determine whether a brain area is injured or uninjured was proposed and its reliability was verified by statistical analyses and CT images.

Quantitative and topographic analysis of the human electroencephalogram (EEG) is a valuable method for evaluating brain function in healthy and diseased states, as those analyses could permit determination of scalp signal sources. For example, it has proven useful methods for mapping the seizure patient’s “epileptogenic zone"1,2. It could also be used to characterize different brain pathological states, such as psychiatric and neurological disorders3. However, whether quantitative and spatial EEG analysis can be used as a technique for mapping injured brain areas has not been sufficiently explored4; recent studies that have attempted to resolve this issue5,6, and the current investigation is based on the conclusions from these reports.

Injured regions in the brain are typically identified using computed tomography (CT) or magnetic resonance imaging (MRI). However, compared with CT or MRI, the EEG method has advantages for its cost, high temporal resolution, flexibility, ease-of-use, performance, and speed As such, using EEG at the bedside can be a powerful tool that provides dynamic information about brain functions7–9, and if applied to the assessment of brain injury, has great theoretical and clinical value10,11.

In this study, an EEG analysis method was developed that is referred to as symmetrical channel EEG signal analysis (SESA). The SESA method is based on hypothetical differences in EEG signal features between injured and uninjured brain areas (applicable for unilateral brain injuries, the most common brain injuries). If the difference is statistically significant, it could be used as an index to judge whether or not a brain area is injured. The SESA method is based on the characteristic anatomical and neurophysiological symmetry and functional offside symmetry of the brain12.

Previous studies have shown that EEG signals for corresponding regions in the cerebral hemispheres have a high degree of similarity in the resting state13,14. If a unilateral brain injury occurs, we can infer that this EEG similarity will decrease and differences will increase in the symmetrical cerebral hemisphere. In the active state, characteristic offside symmetry also increases EEG signal differences15. Indeed, recent developments in brain computer interfaces suggest that pattern recognition of imagined movements are based on the characteristic functional offside symmetry exhibited by the brain16–18. Therefore, it is feasible, both in theory and practice, to map brain injury areas using the SESA method.

We applied signal processing methods and statistical analyses to verify whether changes in EEG signal features (in the symmetrical cerebral hemisphere) are real, and whether they can be used as a method for determining brain injury. A set of criteria (range of ratio values) to distinguish between injured and uninjured brain areas was
also proposed based on SESA. To test whether the proposed criteria were reliable, statistical validation was carried out, and the correspondence between diagnoses made based on SESA criteria and CT images was compared for five selected patients.

**Results**

The SESA method was developed as follows. First, two EEG signal features—approximate entropy (ApEn) and slow-wave coefficient (SWC)—were extracted for all brain channels, and the ratio (Cp value) of signals from injured and uninjured areas for these features was calculated. Statistical analyses of Cp values in population and individual channels were carried out, and a range of ratio values was set as a standard for determining whether a brain area is injured or uninjured. Finally, the reliability of the SESA method was validated using CT images.

**Statistical analysis of Cp values for total injured and uninjured channels.** A statistical analysis was performed to evaluate differences in population Cp values for total injured and uninjured channels. The results are shown in Table 1.

Results from Table 1 indicated that:

1. For ApEn and SWC, in both the resting and active states, a P < 0.01 was observed for population mean Cp values, indicating that injured and uninjured brain channels differed significantly; (2) For the Cp value of ApEn, the value of the injured area during the call name stimulus was much smaller than the value in the resting state;

2. For the Cp value of SWC, the value of the injured area during the call name stimulus was much larger than the value in the resting state.

The Cp value for ApEn during the name call stimulus was smaller than in the resting state. As described in the Methods, in the resting state, the ApEn values for both the injured (e.g., Fp1) and symmetrical uninjured (e.g., Fp2) channels did not change significantly (i.e., Fp1/Fp2 remained constant); however, during the name call stimulus, the ApEn value for Fp2 increased but was unaltered for Fp1 due to injury, resulting in an overall decrease in Fp1/Fp2. Conversely, the Cp value for SWC during the name call stimulus was larger than in the resting state because Fp2 was unchanged while Fp1 increased.

**Statistical analysis of Cp values for individual injured channels.** An analysis of population Cp values for all injured and uninjured channels revealed differences in the Cp values for individual injured channels (Figure 1).

The results showed that:

1. For ApEn and SWC, in both resting and active states, there were statistically significant differences (P < 0.01) for all individual injured channels relative to uninjured channels;

| Features | States  | Mean Cp for injured channels | Mean Cp for uninjured channels | P value |
|----------|---------|------------------------------|-------------------------------|---------|
| ApEn     | Resting | 0.665 ± 0.007                | 0.905 ± 0.009                 | 1.98E-13** |
|          | Name call | 0.584 ± 0.006               | 0.874 ± 0.005                 | 3.05E-12** |
| SWC      | Resting | 1.711 ± 0.047                | 1.343 ± 0.017                 | 5.25E-07**s |
|          | Name call | 1.957 ± 0.166               | 1.517 ± 0.072                 | 9.8E-06** |

*One-way analysis of variance (ANOVA) for control groups showed P values all > 0.05 (not listed in the table), indicating that there were no differences between symmetrical channels for the controls. n = 10 for control and n = 45 for patient groups, respectively.

**P < 0.01 for injured vs. uninjured channels. ApEn, approximate entropy; Cp, ratio of signals from injured and uninjured areas; SWC, slow-wave coefficient.*
(2) The Cp value of ApEn for injured channels during the name call stimulus was smaller than in the resting state; 
(3) The Cp value of SWC for injured channels during the name call stimulus was larger than in the resting state (details in Appendix III).

An explanation for why the Cp value decreases for ApEn while increasing for SWC is given above (see Appendix I).

**Criteria for assessing injured vs. uninjured brain channels.** The above results indicated that there are significant differences in Cp values between injured and uninjured areas as well as individual channels. Thus, statistically determined differences served as a basis for establishing criteria that distinguished between injured and uninjured brain areas. These criteria (section 7 of Methods) and preliminary ranges of Cp values are given in Tables 2 and 3.

**Verification of accuracy.** Permutation testing was carried out to assess the accuracy of the established method. For example, the Cp values of ApEn for Fp1 or Fp2 with the name call stimulus comprised 31 cases; of these, 20 were randomly selected as a training group, and the remaining 11 were the test group. The detailed procedure was as follows: the range of Cp values for the training and test groups were calculated using the above-described method; the reliability was assessed by determining the region of overlap between the Cp values of the training and test groups. This process was repeated for randomly selected training and test cases. After 50 iterations, the average accuracy was found to be 92.3%.

The calculation was repeated for the injured leads of F7 or F8, T3 or T4, and uninjured leads P3 or P4, C3 or C4, F3 or F4, and O1 or O2 (Figure 2). The results showed a high average accuracy, confirming that the method is reliable.

**Calculation of probabilities of type I and II errors.** The probability of occurrence of type I and II errors (incorrectly judging an uninjured area as injured one, or vice versa) was determined. The Cp values of ApEn for Fp1 or Fp2 is used as an example. Of 45 patients, 31 were injured at Fp1 or Fp2, and 14 were uninjured in these locations. Assuming that the 14 cases follow a normal distribution, Cp values of ApEn for Fp1 or Fp2 were calculated as for injured cases. The

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**Table 2 | Range of Cp values for injured channels**

| Features | State       | Fp1 or Fp2 | F7 or F8 | T3 or T4 | T5 or T6 |
|----------|-------------|------------|----------|----------|----------|
| ApEn     | Resting     | 0.583 – 0.699 | 0.626 – 0.744 | 0.593 – 0.753 | 0.629 – 0.717 |
|          | Call Name   | 0.503 – 0.638 | 0.520 – 0.638 | 0.543 – 0.665 | 0.535 – 0.636 |
| SWC      | Resting     | 1.554 – 1.952 | 1.615 – 1.935 | 1.537 – 1.778 | 1.502 – 1.694 |
|          | Call Name   | 1.736 – 2.466 | 1.612 – 2.161 | 1.587 – 2.131 | 1.755 – 2.164 |

A brain area was considered as injured if the corresponding Cp values were within the ranges indicated for F3, F4, P3, P4, C3, C4, O1, and O2 channels. ApEn, approximate entropy; SWC, slow-wave coefficient.

**Table 3 | Range of Cp values for uninjured channels**

| Features | State       | F3 or F4 | C3 or C4 | P3 or P4 | O1 or O2 |
|----------|-------------|----------|----------|----------|----------|
| ApEn     | Resting     | 0.634 – 1.071 | 0.722 – 1.106 | 0.737 – 1.165 | 0.716 – 1.233 |
|          | Call Name   | 0.627 – 1.125 | 0.697 – 1.127 | 0.736 – 1.216 | 0.650 – 1.413 |
| SWC      | Resting     | 0.997 – 1.394 | 1.001 – 1.445 | 1.026 – 1.453 | 0.952 – 1.474 |
|          | Call Name   | 1.056 – 1.545 | 1.111 – 1.709 | 1.146 – 1.611 | 1.118 – 1.651 |

A brain area was considered as uninjured if the corresponding Cp values were within the ranges indicated for F3, F4, P3, P4, C3, C4, O1, and O2 channels. ApEn, approximate entropy; SWC, slow-wave coefficient.

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Figure 2 | Verification of criteria for distinguishing between injured and uninjured brain areas. Data represent the average accuracy after 50 iterations of permutation testing. The lowest average accuracy was from the T5/T6 leads, partly due to a smaller sample size for this group (12 cases).
distribution of the ApEn Cp values for the two groups is shown in Figure 3. The probability of type I and II errors was calculated at the point at which the curves intersect and determined as the areas under the curve.

The calculated probabilities for all leads are shown in Table 4. Since the values were small, it can be concluded that the method is reliable.

**Verification of the SESA method using CT images.** To test the reliability of the proposed criteria, five additional patients were selected for whom diagnoses made based on the SESA method and CT images were compared. The Cp values from one patient are listed in Table 5. In this case, the exact location of the injured area is unknown. Calculating Cp values of the symmetric channel permitted determination of the injured areas. According to the diagnostic criteria, brain areas corresponding to channels with Cp values shown in bold italics in Table 5 (FP1/FP2 and F7/F8) were injured.

An example of results obtained using the SESA method and CT images for one patient is shown in Figure 4. A good correspondence was observed between the two methods.

**Table 4 | Probabilities of type I and II errors**

| Features | Status     | Fp1/Fp2    | F7/F8    | T3/T4    | T5/T6    |
|----------|------------|------------|----------|----------|----------|
|          |            | Type I     | Type II  | Type I   | Type II  | Type I   | Type II  |
| ApEn     | Quiet      | 0.038      | 0.048    | 0.046    | 0.017    | 0.022    | 0.037    | 0.012    | 0.048    |
|          | Call Name  | 0.032      | 0.034    | 0.019    | 0.018    | 0.042    | 0.03     | 0.032    | 0.024    |
| SWC      | Quiet      | 0.021      | 0.019    | 0.039    | 0.022    | 0.023    | 0.043    | 0.045    | 0.030    |
|          | Call Name  | 0.011      | 0.036    | 0.026    | 0.041    | 0.019    | 0.039    | 0.029    | 0.022    |

**Discussion**

The SESA method was developed to map injured brain areas, and its clinical utility was verified using theoretical and experimental methods. Although theoretical simulation studies have been conducted in which functional analyses of asymmetric EEG recordings have been applied towards clinical assessment, our report investigates in which functional analyses of asymmetric EEG recordings have been applied towards clinical assessment. Although theoretical simulation studies have been conducted in which functional analyses of asymmetric EEG recordings have been applied towards clinical assessment, our report investigates in which functional analyses of asymmetric EEG recordings have been applied towards clinical assessment.

**Establishment of diagnostic criteria using patient data.** A range of ratio values defined for each channel is proposed as a standard for determining whether or not a brain area is injured. Although the method was validated by statistical analyses and was proven reliable when applied to the diagnosis of five patients, for routine use in practice (i.e., as an adjuvant clinical diagnostic standard) the accuracy should be confirmed statistically in a larger patient cohort. The present study population included only 45 patients, and it is expected that through cooperative efforts between hospitals, large amounts of data can be obtained that will allow verification of the reliability of this method.

**Selection of EEG signal features.** As many EEG signal features can be used as EEG parameters, the selection of suitable EEG characteristics as parameters is also important for the SESA method. A good EEG parameter represents brain injury states well and has values that can be easily calculated. The ApEn and a defined EEG frequency domain (SWC) were chosen as EEG parameters. Characteristics of nonlinear EEG dynamics include ApEn, sample and permutation entropies, and Lempel-Ziv complexity. Future research will test different features to determine the best for various applications. In terms of EEG frequency domain features, the ratio of the slow wave is a good EEG feature for patients with severe disorders of consciousness. Whether SWC is a good EEG feature for patients with normal consciousness also requires further experimental investigation.

**The spatial resolution of SESA mapping.** We used a 40-channel EEG acquisition instrument to acquire EEG signals, using 16 channels for the actual scalp electrodes. Applying high-density EEG signal recording instruments (with 128 or 256 channels) will greatly increase the spatial resolution of injured areas, facilitating studies that are both clinically relevant and provide basic functional information by revealing the topography of injured brain areas. However, signal interference from the various leads...
Table 5 | Characteristics and results for one patient

| Features  | State    | FP1/FP2 | F7/F8 | T3/T4 | T5/T6 | F3/F4 | C3/C4 | P3/P4 | O1/O2 |
|-----------|----------|---------|-------|-------|-------|-------|-------|-------|-------|
| ApEn      | Resting  | 0.731   | 0.788 | 0.654 | 0.681 | 0.773 | 0.815 | 0.863 | 0.917 |
|           | Call name| 0.702   | 0.691 | 0.624 | 0.586 | 0.786 | 0.773 | 0.843 | 0.902 |
| SWC       | Resting  | 1.274   | 1.491 | 1.657 | 1.674 | 1.164 | 1.311 | 1.403 | 1.197 |
|           | Call name| 1.516   | 1.453 | 1.717 | 1.786 | 1.392 | 1.503 | 1.461 | 1.265 |

EEG acquisition and noise reduction. The EEG acquisition instrument with 40 channels (NuoCheng, Shanghai, China) had 16 channels used to record EEG signals. scalp electrodes were placed according to the international standard 10/20 system and the electrode impedance was ≤5 KΩ, and the sampling frequency was 256 HZ, A1 and A2 were the reference electrodes and Fpz was the grounding electrode.

- **EEG signal acquisition.** EEG signals in two brain states were recorded, one under the resting state and the other under the call-name stimulus state. The two states were recorded to determine which brain state’s EEG signals were better for extracting features for the ratio calculation (further details in the Discussion section).

  For recording in the resting state, 10 min sustainable EEG signals were acquired from each subject. For recording in the call-name stimulus state, the acquisition process is shown in Figure 5. There was a 12 s call-name stimulus recording, during which a selected patient’s family member would call the patient’s name three consecutive times. When the third call ended, data collecting continued for 2 min before ending.

- **EEG noise reduction.** Unstable data caused by external interference (such as EMG interference caused by sudden movements of patient limbs or EEG interference when patients blink during the acquisition process) was manually removed. The toolbox-IIR filter in EEGLAB was processed to remove interference caused by the 50 Hz power frequency.

Extraction of EEG features. Two EEG features were chosen as parameters to compute ratios of symmetrical channels for injured and uninjured brain areas (and channels). One ratio is Approximate Entropy (ApEn), and the other is a defined EEG feature termed the slow wave coefficient (SWC). The extraction processes of the two features are described below.

- **Slow wave coefficient (SWC) calculation.** Eeg rhythm is considered an overall indicator of the excitability of the central nervous system. In general, a lower frequency of slow waves corresponds to a greater degree of abnormality in the corresponding areas, thus, brain damage causes an increase in the slow wave and decrease in the fast wave band in the EEG power spectrum. As such, SWC provides a good index for the identification of damaged brain areas.

  Since the EEG power spectrum is typically divided into six bands, i.e., δ (1.0–4.0 Hz), θ (4.1–8.0 Hz), α (8.1–10.0 Hz), β (10.1–13.0 Hz), β (13.1–17.5 Hz), and β (17.6–30.0 Hz), SWC was defined as:

  \[ SWC = \frac{\text{spectrum of } (\delta + \theta)}{\text{spectrum of } (\alpha + \beta)} \]

  In this formula, \( \alpha = \alpha_1 + \alpha_2, \beta = \beta_1 + \beta_2 \).

  Calculation of SWC was as follows:

  1. 2 min EEG signals of each subject in the resting state and 12 s in call-name stimulus state were intercepted;
  2. The power spectrum of each band was extracted by Fourier transform;
  3. By using the above-mentioned definition of SWC, values of each channel for every subject were obtained.

- **Approximate Entropy (ApEn) calculation.** Approximate Entropy (ApEn) is a nonlinear dynamic character used to describe the complexity and regularity of signals to measure the irregularity of a time series. Its physical significance is the probability of generating a new model of a time series when the dimension changes. The larger probability indicates the sequence is more complex, leading to a greater ApEn. Recent research has shown that nonlinear analysis can directly monitor and measure the suppression level of the cerebral cortex in real-time. For patients with severe disorders of consciousness, the higher entropy indicates the patient is more conscious; thus ApEn is another index that reflects damaged brain areas.

  For the calculation of ApEn a time window was put on each channel of EEG data and 2 s (N = 512) chosen as the window time, then one sampling point was used as a window shift to calculate the ApEn of each channel. After that, the average ApEn from all time windows was taken as the ApEn value. Details of the computing method are provided elsewhere.

Figure 4 | Injured channels determined by the SESA method (left) and CT images of injured area (right) for one patient.
Grouping of symmetrical channels. For the grouping of symmetric channels, the side and area of injury in each patient were identified from CT or MRI images by a clinician. Channels corresponding to injured areas were referred to as injured channels (with the offside channel designated as the symmetric uninjured channel; see Fig. 3). Channels for the whole brain were then divided into injured and uninjured areas (shown as the areas above and below the red line in Fig. 3, respectively), with ratios values of symmetric channels calculated as [injured channels vs. symmetric uninjured channels] and [uninjured channels vs. symmetric uninjured channels], respectively. (as shown the area under the red line in Figure 6).

Computation of ratio values. As the ratio values are critical for the SESA method (according to our proposed hypothesis) the ratio values of EEG features between injured and uninjured brain areas as well as channels were calculated. For APln and SWC, the calculations were as follows: for injured areas, the feature (APln or SWC) of the injured channel was divided by the feature of the symmetric uninjured channel; for uninjured areas, the feature of the uninjured channel was divided by the feature of the symmetric uninjured channel.

Statistical analysis of ratio values in population and individual channels

1. After ratio values were acquired, differences in ratio values for population (i.e., total injured vs. uninjured channels) and individual injured channels (i.e., injured individual F1 vs. total uninjured channels) were evaluated. For convenience, $Cp$ was used instead of ratio values. The analysis of $Cp$ values in population channels was performed as follows:

2. Computation of the mean ratio for all injured area channels;
3. Computation of the mean ratio for all uninjured channels;
4. Placement of 25 means (cases) in a column and one-way ANOVA of the ratio between injured and uninjured channels.

The analysis of $Cp$ values in individual injured channels was performed as follows:

1. The totals for individual injured channels were calculated as 20 FP1 or FP2, 18 F7 or F8, 17 T3 or T4, 5 T5 or T6;
2. The mean ratio for all individual injured channels was calculated;
3. The mean ratio for all individual uninjured channels was calculated;
4. One-way ANOVA was performed for injured and uninjured channel groups.

Figure 5 | Acquisition procedure under the call-name stimulus.

Figure 6 | Diagram of channel grouping. Two kinds of symmetrical channel pairs are shown, corresponding to injured (i.e., injured channel vs. symmetrical uninjured channel; broken dark yellow line) and uninjured (i.e., uninjured channel vs. symmetrical uninjured channel; broken green line) areas.

Establishing criteria for diagnosing an injured area. If the statistical analysis of ratio values revealed significant differences in population and individual channels between injured and uninjured brain areas, a determination of whether a brain injury had occurred was made based on a range of ratio values. SPASS analyses were first used to determine whether the $Cp$ values of all channels follow a normal distribution. By applying statistical theory, preliminary criteria for judgment were derived. For an injured channel (e.g., Fp1 or Fp2), the ratio of 20 patients were taken as sample values to estimate the range of the ratio for corresponding injured brain areas (for other injured channels, the computing methods were the same). For an uninjured channel (e.g., F3–F4, C3–C4, P3–P4, O1–O2), the ratios of 25 patients were regarded as sample values to estimate the range of the ratio for corresponding uninjured brain areas. More details about the computing methods are in Appendix III.
21. Athina, T. The timing of exploratory decision-making revealed by single-trial topographic EEG analyses. *NeuroImage* **60**, 1959–69 (2012).
22. van Putten Michel, J. A. M. *et al.* A brain symmetry index (BSI) for online EEG monitoring in carotid endarterectomy. *Clin. Neurophysiol.* **115**, 2157–2168 (2011).
23. Abbas, B.-F. & Hamid, S.-Z. Multi-area neural mass modeling of EEG and MEG signals. *NeuroImage* **52**, 793–811 (2010).
24. Husain, A. M. Electroencephalographic assessment of coma. *J. Clin. Neurophysiol.* **23**, 208–220 (2006).
25. Andrew, M. G., Jonathan, D. V., Mary, M. C., Jonathan, C. B. & nNicholas, D. S. Determination of awareness in patients with severe brain injury using EEG power spectral analysis. *Clin.Neurophysiol.* **11**, 2159–2168 (2011).
26. Abisolo, D. *et al.* Entropy analysis of the EEG background activity in Alzheimer’s disease patients. *Physiol. Meas.* **27**, 241 (2006).
27. Ranannd, P., Nampoori, V. P. N. & Sreenivasan, R. Complexity quantification of dense array EEG using sample entropy analysis. *J. Integr. Neurosci.* **3**, 343–358 (2004).
28. Abisolo, D. *et al.* Analysis of EEG background activity in Alzheimer’s disease patients with Lempel–Ziv complexity and central tendency measure. *Med. Eng. Phys.* **28**, 315–322 (2006).
29. Kotchoubey, B. *et al.* Information processing in severe disorders of consciousness: vegetative state and minimally conscious state. *Clin. Neurophysiol.* **116**, 2441–2453 (2005).
30. Gosseryes, O., Schnakers, C., Ledoux, D. *et al.* Automated EEG entropy measurements in coma, vegetative state/unresponsive wakefulness syndrome and minimally conscious state. *Funct. Neurol.* **26**, 25–30 (2011).
31. Okamoto, M. *et al.* Three-dimensional probabilistic anatomical cranio-cerebral correlation via the international 10–20 system oriented for transcranial functional brain mapping. *NeuroImage* **21**, 99–111 (2004).
32. Herwig, U., Satrapi, P. & Schönfeldt-Lecuona, C. Using the international 10–20 EEG system for positioning of transcranial magnetic stimulation. *Brain Topogr.* **16**, 95–99 (2003).
33. Di, H. B., Yu, S. M., Weng, X. C. *et al.* Cerebral response to patient’s own name in the vegetative and minimally conscious states. *Neurology* **68**, 895–899 (2007).
34. Pincus, S. Approximate entropy (ApEn) as a complexity measure. *Chaos* **5**, 110–117 (1995).
35. Adler, G. & Brassen, S. Short-term rivastigmine treatment reduces EEG slow-wave power in Alzheimer patients. *Neuropsychobiology* **43**, 273–276 (2001).
36. Leon-Carrion, J. *et al.* Brain function in the minimally conscious state: a quantitative neurophysiological study. *Clin. Neurophysiol.* **119**, 1506–1514 (2008).
37. Richman, J. S. & Moorman, J. R. Physiological time-series analysis using approximate entropy and sample entropy. *AM. J. Physiol-Heart C.* **278**, H2039–H2049 (2000).
38. Pincus, S. M. Approximate entropy as a measure of system complexity. *P. Natl. Acad. Sci.* **88**, 2297–2301 (1991).

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**Author contributions**

Y.Y. and L.Y. conceived the project and experiment designed; L.Y. and L.X.P. wrote the manuscript draft; Y.Y., L.X.P. and L.L.H. co-revise the manuscript; L.X.P., Z.D.K. and Y.W.W. processed data and prepared the figures; L.J.Q. and L.X.P. collected patients. All authors reviewed the manuscript.

**Additional information**

**Supplementary information** accompanies this paper at http://www.nature.com/scientificreports

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