Chronobiological patterns of acute aortic dissection in central China

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ABSTRACT

Background Acute aortic dissection (AAD) is a life-threatening emergency with poor clinical outcomes. Understanding the chronobiological patterns of AAD onset would be helpful for identifying the triggers of AAD and preventing this catastrophic event.

Methods We collected data from 2048 patients diagnosed with AAD at Tongji Hospital (Wuhan, China) from 2011 to 2018. The χ² test was used to determine whether a specific period had significantly different seasonal/weekly distributions from other periods. Fourier models were used to analyse the rhythmicity in monthly/circadian distribution.

Results The mean age was 53.4±10.9 years, and 1161 patients (56.7%) were under 55 years. One thousand six hundred fifty-seven patients (80.9%) were male, and 935 cases (45.7%) were type A dissections. The proportions of patients with comorbid hypertension/diabetes were 60.3% (1234 cases) and 1.8% (36 cases), respectively. A peak was identified in colder periods (winter/December) and a trough in warmer periods (summer/June). No significant variation was observed in weekly distribution. Fourier analysis showed a statistically significant circadian variation (p<0.001) with a nocturnal trough in 2:00–3:00, a morning peak in 9:00–10:00, and an afternoon peak in 16:00–17:00. Subgroup analyses identified circadian rhythmicity in all subgroups except for the female group and younger group (younger than 55 years).

Conclusion Our results confirmed that the onset of AAD exhibits significant seasonal, monthly, and circadian patterns. Patients with AAD with different Stanford-type dissections, sexes, ages and hypertension statuses could present different circadian variations. These findings may provide novel perspectives for identifying the triggers of AAD and better preventing this catastrophic event.

INTRODUCTION

Acute aortic dissection (AAD) is a life-threatening emergency with high morbidity and mortality.1 The reported incidence of AAD is three to five cases per 100 000 people every year worldwide,2 but this incidence might be underestimated since hospital-based reports did not take predismension deaths into account. A study based on a large cardiovascular registry indicated that more than 60% of AAD cases were not detected clinically but were first identified by autopsy.3 Therefore, despite the great advancements in therapeutic techniques, the early recognition and diagnosis of AAD are still of vital importance. Epidemiological evidence suggests that the onset of AAD occurs at different times with different probabilities.4 Analysing the chronobiological patterns of AAD could contribute to the prevention and timely treatment of this condition.

Currently, compelling evidence has revealed that myocardial infarction, stroke and arrhythmias occur more frequently in the morning hours, in winter and on Mondays.3–7 Disturbances to the internal circadian clock, modifications of behavioural patterns and fluctuations of environmental variables may contribute to the increased risks at specific times. However, limited studies have discussed how the time of day, week and year affect the onset of AAD, and most of these studies were conducted in Western countries with limited sample sizes and without subgroup analyses.8–13 To evaluate the effect of chronobiological patterns on the occurrence of AAD, we aimed to expound the temporal trends of AAD onset in an 8-year time-series study in central China, and we believe that these data could help predict and prevent the disastrous outcomes of this condition.

METHODS

Study population and data collection

This time-series study was reported according to the Strengthening the Reporting of Observational Studies in Epidemiology checklist. This study was based on electronic medical records of patients admitted to Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology, which is a major comprehensive hospital in central China that mainly treats patients from Hubei Province. We enrolled all patients diagnosed with AAD from 1 January 2011 to 31 December 2018 based on their electronic medical records. International Classification of Diseases-10 codes (I71.000–I71.005, I71.007) were used to preliminarily identify patients with aortic dissection (AD). Imaging information (ie, CT angiography and transoesophageal echocardiography) and surgical records of the included patients were checked for further confirmation. Since the clinical manifestations of AAD, intramural haematoma (IMH) and penetrating atherosclerotic ulcer (PAU) were similar and a clear diagnosis among the three was challenging, we included only patients who had imaging evidence or surgical records supporting the diagnosis of AAD. To analyse the time distribution of AAD onset, we set predefined exclusion criteria for the participants to improve the quality and validity of our research. The exclusion criteria were as follows: (1) AD without typical symptoms, defined as acute chest/back/abdominal pain within 14 days,4 since the onset time could not be

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accurately recalled; (2) simple IMH/simple PAU; (3) AD with congenital abnormalities, such as Marfan syndrome (MFS) or bicuspid aortic valve or a history of cardiovascular surgery/atherogenic AD; (4) traumatic AD/AD in pregnancy; and (5) aortic aneurysm (since only dissections were considered for inclusion). The following baseline characteristics of the included patients were extracted separately: sex, age, Stanford type of AD, and medical history of diabetes and hypertension. The onset date and exact time (if available) of AAD were inferred from the occurrence of acute symptoms based on electronic medical records.

Study design
We aimed to investigate four temporal patterns of AAD onset, including seasonal, monthly, weekly and circadian distributions. The included patients were categorised into 4 seasonal periods, 12 1-month periods and 7 1-day periods. Four seasons were defined according to the local climate conditions: spring from March to May, summer from June to August, autumn from September to November, and winter from December to February. For patients with a precise onset time, circadian rhythmicity was analysed after categorisation into 24 1-hour increments (eg, onset time between 0:00 and 0:59 categorised in the 0:00 group). Furthermore, we investigated whether differences in sex, age (younger than 55 years old, older or equal than 55 years old), AAD type (Stanford classification) and comorbid conditions such as hypertension would affect the chronobiology pattern, especially circadian variations.

Statistical analysis
Continuous variables following a normal distribution are described as the mean±SD, while those that were non-normally distributed are described as the median (IQR). Categorical variables are described as percentages. The seasonal and weekly distributions of AAD onset were tested for uniformity among the total population and subgroups by the $\chi^2$ test. Values following uniform distribution were used as expected values to complete the test. For the monthly data, a Fourier model with one harmonic of 12 months was utilised to help analyse the trend and rhythmicity. For the circadian data, a Fourier model with four harmonics (periods of 24, 12, 8 and 6 hours) was chosen to fit the data based on the preliminary test. R squared was used to represent the goodness of fit. The F statistic (hypothesis of straight-line model fitting) was reported to represent the statistical significance of the fitted function. A value of $p<0.05$ was considered significant in all tests. All statistical analyses were performed on GraphPad Prism V8.3.0.

Patient and public involvement statement
This planned study was not presented to a patient group for their input prior to commencement of the study due to its retrospective nature. When we extracted data from the electronic medical records, we first deleted all personal information, including names and hospitalisation numbers, to ensure that the privacy of the patients would not be disclosed to the researchers and other involved members.

RESULTS
Basic characteristics of the study population
In total, 2048 patients with AAD with typical symptoms and definitive diagnoses were identified (see figure 1), including 935 Stanford type A dissections (45.7%) and 1113 Stanford type B dissections (54.3%). One thousand six hundred fifty-seven patients (80.9%) were male. The average age was 53.4±10.9 years, and 1161 patients (56.7%) were younger than 55 years old. The proportions of patients comorbid with hypertension and diabetes were 60.3% and 1.8%, respectively. Among the included patients, 1517 patients had a precise onset time of AAD in their medical records, and we included these patients in the circadian analysis. Details are shown in table 1.

Seasonal, monthly and weekly variations
The seasonal distribution of AAD showed significant differences ($\chi^2$ 93.97, df 3, $p<0.0001$) (see figure 2). In general, most cases occurred in winter (34.2%, 701 cases), and the fewest occurred in summer (14.8%, 303 cases). This trend also existed in the group of Stanford type B dissections ($\chi^2$ 45.81, df 3, $p<0.0001$), with 399 cases (35.8%) occurring in winter and 176 cases (15.8%) occurring in summer. For Stanford type A dissections ($\chi^2$ 55.12, df 3, $p<0.0001$), the fewest cases also occurred early in the season.
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in summer (13.7%, 127 cases), and the number of cases that occurred in autumn (33.4%, 312 cases) was almost equal to that in winter (32.3%, 302 cases).

Fourier rhythm analysis identified a statistically significant temporal pattern ($p<0.001$) in monthly distribution (see figure 3). A peak was observed around December, and a nadir was observed around June. Similar patterns were detected in patients with type A and type B dissections.

The weekly distribution of AAD is displayed in figure 4. No statistically significant difference was observed in the number of cases on different days of the week ($\chi^2 2.439$, df 6, $p=0.875$), neither in patients with type A dissection ($\chi^2 2.998$, df 6, $p=0.809$) nor in patients with type B dissection ($\chi^2 5.311$, df 6, $p=0.505$).

Circadian variations

Fourier rhythm analysis identified statistically significant circadian patterns among the total population ($p<0.001$), with a nocturnal trough between 2:00 and 3:00, a minor morning peak between 9:00 and 10:00, and a major afternoon peak between 16:00 and 17:00 (see figure 5).

Similar trends were detected in all subgroups except for the female group ($p=0.164$) and the younger group ($p=0.080$). The nocturnal trough appeared between 2:00 and 4:00 in all groups...
except in the female group (at 4:58). Most subgroups showed a morning peak between 8:00 and 10:00, except for an earlier peak for the hypertensive group (at 7:00) and a delayed peak for the older group (at 10:40). The afternoon peak occurred between 16:00 and 17:00 in all subgroups except the female group, which had a peak at approximately 18:29. Detailed information is displayed in table 2 and figure 6.

DISCUSSION
We conducted a single-centre retrospective study including 2048 patients from central China to identify potential chronobiological patterns of AAD onset. First, we recognised a higher incidence in the colder period, such as in winter or December, and a lower incidence in the warmer period, such as in summer or June. These results are consistent with most previous studies, and the winter peak was reported to be independent of climate conditions. Lower temperature is associated with increased sympathetic activity and blood pressure (BP), as well as changes in haematological properties (including increased blood viscosity, haemoglobin, mean corpuscular volume, platelet aggregability, factor VII, fibrinogen, and acute-phase reactants and decreased red cell deformability). Moreover, the concentrations of air pollutants, especially fine particulate matter, are also elevated in winter and are associated with aortic plaque progression, hypertension and aortic constriction. These changes may lead to a surge in the shear forces exerted on the aortic intima, triggering AAD in colder months.

Second, no significant weekly variation was observed for AAD onset in our study. The results from previous studies on the weekly distribution of AAD onset remain conflicting. Some studies reported a Monday peak, while others reported a Wednesday peak. The weekly distribution of cardiovascular disease (CVD) may reflect physiological fluctuations resulting from an increase in physical and mental burdens from leisurely weekend activities to stressful weekdays. Different working statuses and occasional holidays may easily alter the weekly patterns of AAD onset, leading to a non-significant result.

Moreover, we demonstrated a significant circadian pattern in AAD onset. For the entire studied population, a nocturnal trough appeared at 2:52, followed by a minor morning peak at 9:27 and a major afternoon peak at 16:41. Indeed, a similar daily pattern has been found in the occurrence of many adverse cardiovascular events, including stroke, myocardial infarction and sudden cardiac death, which all occur most frequently in the morning. Such a circadian pattern may result from fluctuations in BP. It has been known for several decades that BP exhibits a circadian rhythm in humans due to an innate circadian rhythm, as well as external stimulation from environmental fluctuations and daily behaviours—BP dips at night during rest, undergoes a steep increase in the morning (known as the 'morning surge')

Table 2 Circadian variations of hourly data of AAD onsets, fitted by Fourier model

| Group          | N   | Nocturnal trough | Morning peak     | Afternoon peak   | R squared | P value |
|----------------|-----|------------------|------------------|------------------|-----------|---------|
| Total          | 1517| 2:52             | 37.82±11.74      | 9:27             | 16:41     | 0.8151  | 0.0008  |
| Type A         | 737 | 2:36             | 15.43±7.618      | 8:29             | 16:41     | 0.4497  | 0.7203  |
| Type B         | 780 | 4:03             | 21.30±8.914      | 10:00            | 16:41     | 0.5197  | 0.0143  |
| Male           | 1242| 2:50             | 29.23±9.248      | 9:34             | 16:39     | 0.7922  | 0.0004  |
| Female         | 275 | 4:58             | 7.06±5.282       | 8:46             | 18:29     | 0.7390  | 0.1641  |
| Age <55 years  | 875 | 4:07             | 23.32±10.628     | 8:54             | 16:51     | 0.5287  | 0.0803  |
| Age ≥55 years  | 642 | 2:18             | 11.38±4.919      | 10:40            | 16:34     | 0.4050  | <0.0001 |
| Hypertensive   | 929 | 3:02             | 23.86±9.297      | 7:00             | 16:50     | 0.6188  | 0.0088  |
| Normotensive   | 577 | 1:57             | 13.55±4.834      | 9:32             | 16:14     | 0.3423  | 0.0002  |

Letter N indicates the number of cases in each group; letter n shows the number of cases fitted by Fourier model, presented by numbers±95% CIs; R square represents goodness of the model; and P, probability value resulting from the F statistic used to test hypothesis of straight-line model fitting. AAD, acute aortic dissection.

Figure 6 Circadian variations of AAD onset for different subgroups using a Fourier model. Each point represents the total count of onset AAD for the periods considered. The active line is the overall best-fitting curve calculated by rhythm analysis, resulting from four significant harmonics with periods of 24, 12, 8 and 6 hours. The filled area represents 95% CI. AAD, acute aortic dissection.
and peaks typically in the late afternoon. Moreover, the reactivity of BP to daily physical activity has been reported to show 24-hour variations, in which systolic BP reactivity was highest in the morning and exhibited a secondary rise in the afternoon. Most cardiovascular-related variables exhibit a circadian rhythm with a morning peak, including heart rate, operation of the renin–angiotensin–aldosterone system, vascular resistance and sympathetic activity. In addition, the balance between coagulation and fibrinolysis is also disturbed, resulting in hypercoagulability and hypofibrinolysis in the morning. The surges in BP and BP reactivity, together with other changes in the cardiovascular system, would increase arterial shear force, triggering the onset of AAD in corresponding periods, especially in people with a genetic predisposition.

To further investigate the temporal pattern, we performed circadian analyses in subpopulations and identified some intriguing nuances. Patients with type A dissection showed a higher morning surge compared with patients with type B dissection. Type A dissection is closely related to inherited connective tissue diseases, while type B dissection is more likely to be related to atherosclerosis. However, previous studies on a genetic susceptible model (patients with MFS) and an atherosclerotic model (patients with spontaneous rupture of abdominal aortic aneurysm) reported similar circadian patterns. Whether patients with different types of AAD present different circadian patterns needs further investigation. Moreover, female patients did not exhibit a significant circadian pattern. Women with AAD have different clinical features and were reported to have a temporal delay from symptom onset to diagnosis, which may partly explain the non-significant result. The smaller sample size can also affect the model fitting process, leading to the non-significant result. In addition, the normotensive group exhibited a more obvious morning peak than the hypertensive group. Notably, abnormalities or loss of circadian rhythmicity in BP is common in certain forms of arterial hypertension, which may change the original onset patterns of AAD. Treatment for hypertension may also alter the innate circadian pattern of BP. However, whether treatment for hypertension or hypertension itself leads to the atypical morning peak needs further investigation. We also found that the younger patients exhibited a later nocturnal trough (at 4:07) than the older patients (at 2:18), which might be partially due to the different sleeping and waking habits between young and old patients.

A number of epidemiological studies have explored circadian patterns of AAD onset. In 2002, Mehta et al conducted a study that included 957 patients with AAD from around the world enrolled in the International Registry of Aortic Dissection and used non-linear Fourier analysis for the first time. They reported a nocturnal trough, a major early morning peak and a minor afternoon peak. Vitale et al performed a meta-analysis of eight studies, analysing hourly variations using the data from 1816 patients. They reported an increased risk of acute aortic rupture or dissection between 6:00 and 12:00 compared with the risk in other hours. Takagi et al analysed data from seven studies with 1827 patients with AAD, and they identified a peak in 8:00–10:00 and a nadir in 0:00–2:00. In this study, we identified a higher afternoon peak among Chinese patients with AAD, and some Asian studies reported similar results. This nuance could result from different social and dietary habits in different cultures. Asians tend to consume strong liquor and salty food at dinner in the late afternoon, which might account for the elevation in BP and contributes to the higher afternoon peak of AAD onset. We expect more studies on various ethnicities to elucidate this difference.

In summary, we thoroughly explored the seasonal, monthly, weekly and circadian patterns of AAD onset in an 8-year time-series study in central China. Understanding the chronological patterns of AAD onset could be helpful for preventing this catastrophic event. In addition, we performed circadian analyses among subgroups and identified some intriguing nuances. These findings may provide novel perspectives for clinicians to understand the triggers of AAD. Moreover, with the identified circadian patterns of AAD onset, we should place more emphasis on chronotherapy for AAD. The intake of antihypertension or anticoagulation drugs in the evening could better match the daily physiological oscillations and significantly reduce CVD morbidity and mortality. These factors could probably benefit people with risk factors for AD, such as hypertension or atherosclerosis.

There are several limitations in this study. First, our study had a single-centre retrospective design despite the large study population. There could be inclusion bias from using data extracted from electronic medical records, and part of the data were based on patient recall. Second, given that AAD has a high mortality rate within hours of onset, patients might fail to reach a hospital for timely medical treatment. Failure to include these patients inevitably affects the accuracy of the results.

**CONCLUSION**

In conclusion, we used data from 2048 Chinese patients and confirmed that the onset of AAD exhibits significant circannual and circadian patterns. For circannual variations, a peak was identified in colder periods, and a trough was identified in warmer periods. For daily variations, we identified a nocturnal trough, a morning peak and an afternoon peak. Differences in AD type, sex, age and hypertension status could lead to different circadian variations. These findings may provide novel perspectives to help us identify the triggers of AAD and better prevent this catastrophic event.

**Key messages**

What is already known on this subject?

- Epidemiological evidence suggests that the onset of AAD exhibits significant circannual and seasonal/monthly variations. However, most of these studies were conducted in Western countries and lack subgroup analyses.

What might this study add?

- This 8-year time-series study used data from 2048 patients in central China and identified significant seasonal, monthly and circadian patterns of AAD onset. More AAD cases occurred in colder periods (winter/December), and fewer cases occurred in warmer periods (summer/June). Rhythm analysis indicated a nocturnal trough, a morning surge and an afternoon peak in circadian patterns. Patients with different AAD types, sexes, ages and hypertension statuses presented different circadian variations.

How might this impact on clinical practice?

- Understanding the chronological patterns of AAD onset will help clinicians identify the triggers of AAD and contribute to the prevention and timely treatment of this condition.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were involved in the design, conduct, reporting or dissemination plans of this research. Refer to the Methods section for further details.

Patient consent for publication Not required.

Ethics approval This study was approved by the Research Ethics Commission of Tongji Medical College (2019-S1149). Informed consent form was not applicable for this study.

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Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

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