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Chest Pain Center Accreditation Is Associated With Improved In-Hospital Outcomes of Acute Myocardial Infarction Patients in China: Findings From the CCC-ACS Project

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Background—Chest pain center (CPC) accreditation plays an important role in the management of acute myocardial infarction (AMI). However, no evidence shows whether the outcomes of AMI patients are improved with CPC accreditation in China.

Methods and Results—This retrospective analysis is based on a predesigned nationwide registry, CCC-ACS (Improving Care for Cardiovascular Disease in China-Acute Coronary Syndrome). The primary outcome was major adverse cardiovascular events (MACE), including all-cause death, reinfarction, stent thrombosis, stroke, and heart failure. A total of 15 344 AMI patients, from 40 CPC-accredited hospitals, were enrolled, including 7544 admitted before and 7800 after accreditation. In propensity score matching, 6700 patients in each group were matched. The incidence of 7-day MACE (6.7% versus 8.0%; \( P = 0.003 \)) and all-cause death (1.1% versus 1.6%; \( P = 0.021 \)) was lower after accreditation. In multivariate adjusted mixed-effects Cox proportional hazards models, CPC accreditation was associated with significantly decreased risk of MACE (hazard ratio: 0.78; 95% CI, 0.68–0.91) and all-cause death (hazard ratio: 0.71; 95% CI, 0.51–0.99). The risk of MACE and all-cause death both followed a reverse J-shaped trend: the risk of MACE and all-cause death decreased gradually after achieving CPC accreditation, with minimal risk occurring in the first year, but increased in the second year and after.

Conclusions—Based on a large-scale national registry data set, CPC accreditation was associated with better in-hospital outcomes for AMI patients. However, the benefits seemed to attenuate over time, and reaccreditation may be essential for maintaining AMI care quality and outcomes. (J Am Heart Assoc. 2019;8:e013384. DOI: 10.1161/JAHA.119.013384.)

Key Words: accreditation • acute myocardial infarction • chest pain center • China • in-hospital outcomes

Since 1981, when the concept of the chest pain center (CPC) was introduced, studies have shown that CPC is associated with improving timing of chest pain diagnosis, reperfusion time of ST-segment-elevation myocardial infarction (STEMI) and reduced readmissions and costs.1–4 Accreditation by the Society of Chest Pain Centers (now known as the American College of Cardiology [ACC] accreditation committee) has further standardized CPCs in the United States, and the ACC accreditation committee has promoted international accreditation standards since 2010.5–7 Several countries have also established their national CPC accreditation projects, such as the Chest Pain Unit Certification Working Group issued by the German Society of Cardiology.8–10

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Accompanying Appendix S1, Data S1, Tables S1 through S3 and Figures S1 through S6 are available at https://www.ahajournals.org/doi/suppl/10.1161/JAHA.119.013384

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Clinical Perspective

What Is New?

- This is the first study focus on the association of chest pain center accreditation with improved in-hospital major adverse cardiovascular events of acute myocardial infarction patients in China based on a large-scale national registry covering 15 344 patients from 40 hospitals.

What are the Clinical Implications?

- This study demonstrated and confirmed the importance and effectiveness of chest pain center development and accreditation in China.
- Hazards for the risk of outcomes during the in-hospital period as a function of the duration of accreditation followed a reverse J-shaped trend, with the maximum associated effectiveness occurring up until the first year after accreditation, supporting the rationale of the recently established CPC reaccreditation project to ensure a long-lasting effect after initial CPC accreditation.

The incidence and mortality of acute myocardial infarction (AMI) are increasing in China, and the in-hospital mortality of STEMI patients has not improved over the past decade.\textsuperscript{11–13} The development of CPCs in China was initiated in 2010, and the headquarters of China Chest Pain Centers, which oversees CPC accreditation, was officially established in July 2016 to coordinate social resources and promote the rapid development of CPCs.\textsuperscript{14,15} To date, evidence has not shown whether the outcomes of AMI patients are improved by CPC accreditation in China. The purpose of this study was to evaluate whether patients admitted for AMI after CPC accreditation had better in-hospital outcomes than those admitted before accreditation and to further study the temporal associations of the accreditation process on AMI quality improvement.

Methods

For the concern about intellectual property and patient privacy, the data, analytic methods, and study materials will not be made available to other researchers for purposes of reproducing the results or replicating the procedure.

Study Design

The CCC-ACS (Improving Care for Cardiovascular Disease in China–Acute Coronary Syndrome) project is a nationwide registry and quality improvement study with an ongoing database focusing on quality of acute coronary syndrome care. Details of the study design and methodology of the CCC-ACS project have been described elsewhere.\textsuperscript{16} A standard procedure was used during data collecting from the patients’ medical records, and third-party research associates performed regular quality audits to ensure the accuracy and completeness of research data. Institutional review board approval was granted for this research by the ethics committee of Beijing Anzhen Hospital, Capital Medical University, and no informed consent was required.

Study Population

From November 1, 2014, to June 30, 2017, a total of 63 641 patients diagnosed with acute coronary syndrome from 150 hospitals were registered in the database, among which 19 270 patients from 40 accredited hospitals during this period were included in this study. We excluded 2659 patients who were admitted to 10 hospitals with data only before or after accreditation and 1182 patients diagnosed with unstable angina pectoris. In addition, 85 patients with missing data about key variables for the identification of in-hospital outcomes and candidate factors were excluded. Finally, 15 344 patients were included in the final analyses (Figure 1).

In-Hospital Outcomes

In this study, the primary outcome was major adverse cardiovascular events (MACE), including all-cause death, reinfarction, stent thrombosis, stroke, and heart failure during hospitalization. The secondary outcome was in-hospital all-cause death. Only those incidents that occurred within 7 days

Figure 1. Flow diagram of selection of the study population. CCC-ACS indicates Improving Care for Cardiovascular Disease in China-Acute Coronary Syndrome; UAP, unstable angina pectoris.
of admission were taken into account for the analysis because the main effect of the CPC was reflected a short time after the first medical contact; there will be more censored data because of hospital discharge.

Statistical Analysis

Hospital information, demographic characteristics, medical history, clinical procedures, and 7-day in-hospital outcomes of the participants were compared by hospital status of accreditation (before and after) when each patient was admitted. Continuous variables were shown as mean±SD. Categorical variables were presented as the number (percentage). Differences in various characteristics among the groups were compared using the Student t test and χ² test.

The differences of 7-day in-hospital outcomes between the 2 groups were compared in a propensity score–matched population to minimize selection bias from the real world. Patients admitted to hospitals before accreditation were matched at a 1:1 ratio with randomly selected patients admitted to hospitals after accreditation, on the basis of nearest neighbor in terms of Mahalanobis distance with a caliper of 0.02. Propensity score was estimated with a logistic regression model with the variables of age; sex; hospital location (first-line municipality, provincial capital, or prefecture-level city); first medical contact site (no or yes); department arrived (emergency and catheter lab, or outpatient and others); comorbidities including smoking status, diabetes mellitus, hypertension, and dyslipidemia; heart failure history; renal failure history; previous myocardial infarction (MI); previous percutaneous coronary intervention or coronary artery bypass grafting; type of MI; Killip classes; preadmission use of aspirin; preadmission use of P2Y₁₂ receptor inhibitors; and preadmission use of statins.

The incidence of 7-day in-hospital outcomes by hospital status of accreditation in the whole study population and in the propensity score–matched subset were compared. Because some variables were not comparable between the 2 groups even after propensity score matching, we used mixed-effects Cox proportional hazards models containing Gaussian random effects, also known as frailty models, to compare the risk by adjusting other confounders in the logistic regression model, considering that this data set contained hospital-level information and the data structure was hierarchical.

Subgroup analyses of 7-day in-hospital outcomes were then performed according to important characteristics, including age (<65 or ≥65 years), sex (male or female), city type (provincial capital or prefecture-level city), first medical contact site (no or yes), department arrived (emergency and catheter lab, or outpatient and others), type of MI (STEMI or non-STEMI), Killip class I (no or yes), previous MI (no or yes), and previous percutaneous coronary intervention or coronary artery bypass grafting (no or yes).

Considering the quality improvement effect of the CCC-ACS project itself along with time, we performed sensitivity analyses in patients admitted to hospitals without accreditations during the project and those contemporarily admitted in hospitals after accreditation. Propensity score was also estimated with a logistic regression model with all of the listed variables included plus the admission date. Cox proportional hazards models were then used to compare the risk of 7-day in-hospital outcomes.

A 2-tailed P<0.05 was considered statistically significant in all analyses. R software (http://www.R-project.org) was used for all statistical analyses.

Results

Patient Characteristics

A total of 15 344 patients were enrolled in this study from 40 hospitals, including 7544 admitted before the accreditation and 7800 after accreditation. Baseline characteristics are shown in Table 1. Compared with patients admitted to hospitals when not accredited, those admitted after accreditation were mainly from hospitals in cities of provincial capitals, had the hospital as the first medical facility, and arrived in the emergency department and catheter lab. The postaccreditation group had more patients diagnosed with STEMI and had a higher proportion of patients with Killip class I. Comparing comorbidities, patients in the postaccreditation group were less likely to have hypertension, dyslipidemia, heart failure history, and previous MI but more likely to have previous percutaneous coronary intervention or coronary artery bypass grafting. In addition, there were fewer patients to be treated with statins in the postaccreditation group regarding preadmission medications.

After propensity score matching, postmatching absolute standardized differences were <10% for all covariates. The mirrored histograms before and after matching are shown in Figure S1 to present the propensity score distribution status. A total of 6700 patients in each group were matched. The characteristics of the 2 groups were recompared. In the propensity score–matched population, there were no significant differences of baseline characteristics between the 2 groups except for hypertension comorbidity (Table 1).

Seven-Day In-Hospital Outcomes

In-hospital outcomes within 7 days of admission were compared according to accreditation status (Table 2). In the whole study population, the incidence of MACE was significantly lower in patients admitted after accreditation compared with those admitted before accreditation (6.4% versus 8.4%; P<0.001), mainly due to heart failure. The incidence of all-cause death was
also significantly lower after versus before accreditation (1.1% versus 1.6%; \( P = 0.016 \)). Cumulative hazards of MACE and all-cause death were also lower in the postaccreditation group compared with the preaccreditation group (log rank \( P < 0.001 \) and \( P = 0.020 \), respectively; Figure 2A and 2C).

After propensity score matching, the incidence of 7-day in-hospital MACE (6.7% versus 8.0%; \( P = 0.003 \)) and all-cause death (1.1% versus 1.6%; \( P = 0.021 \)) was still lower in the postaccreditation group (log rank \( P = 0.003 \) and \( P = 0.020 \), respectively; Table 2, Figure 2B and 2D).

In mixed-effects Cox proportional hazards models, CPC accreditation was associated with statistically significantly decreased risk of MACE (hazard ratio [HR]: 0.78; 95% CI, 0.68–0.91; \( P = 0.001 \)) and all-cause death (HR: 0.71; 95% CI, 0.51–0.99; \( P = 0.042 \)) after multivariate analysis adjusted for possible confounders (Table 3).

Hazards for the risk of outcomes during the in-hospital period by the duration of accreditation in the propensity score–matched population are presented in Figure 3. The risk of MACE and all-cause death for those patients admitted after the accreditation both follow a reverse J-shaped trend: the risk of both MACE and all-cause death decreases gradually with CPC accreditation, with the minimal risk occurring in patients admitted from 6 months to the first year after accreditation, and then risk increases in the second year and after.

### Subgroup Analyses

Subgroup analyses for MACE and all-cause death based on baseline information were performed in the propensity score–matched population (Figures S2 and S3). The main results were not significantly changed in most subgroups. No interactions

### Table 1. Baseline Characteristics of the Study Population

| Variable                        | Unmatched Before Accreditation | After Accreditation | Propensity Score Matched Before Accreditation | After Accreditation | \( P \) Value |
|---------------------------------|--------------------------------|--------------------|-----------------------------------------------|--------------------|--------------|
| \( n \)                          | 7544                           | 7800               | 6700                                          | 6700               |              |
| Age, y                           | 62.48 (12.61)                  | 62.52 (12.52)      | 0.838                                         | 62.36 (12.62)      | 0.782        |
| Female sex                       | 1796 (23.8)                    | 1808 (23.2)        | 0.369                                         | 1564 (23.3)        | 0.984        |
| Hospital location: provincial capital | 4554 (60.4)                | 5122 (65.7)        | <0.001                                        | 4240 (63.3)        | 0.281        |
| First medical facility: hospital | 3712 (49.2)                    | 4124 (52.9)        | <0.001                                        | 3367 (50.3)        | 1.000        |
| Department: Emergency/catheter lab | 5278 (70.0)                   | 6046 (77.5)        | <0.001                                        | 5025 (75.0)        | 0.564        |
| Type of MI                        |                                |                    |                                               |                    |              |
| STEMI                            | 5296 (70.2)                    | 5659 (72.6)        | 0.001                                         | 4824 (72.0)        | 0.908        |
| Killip class                     |                                |                    |                                               |                    |              |
| I                                | 5298 (70.2)                    | 5797 (74.3)        | <0.001                                        | 4835 (72.2)        | 0.651        |
| II to III                       | 1966 (26.1)                    | 1735 (22.2)        | 1624 (24.2)                                   | 1593 (23.8)        |              |
| IV                               | 280 (3.7)                      | 268 (3.4)          | 241 (3.6)                                     | 228 (3.4)          |              |
| Comorbidity                     |                                |                    |                                               |                    |              |
| Current smoking                 | 3577 (47.4)                    | 3760 (48.2)        | 0.335                                         | 3213 (48.0)        | 0.809        |
| Hypertension                    | 4052 (53.7)                    | 3992 (51.2)        | 0.002                                         | 3534 (52.7)        | 0.013        |
| Dyslipidemia                    | 543 (7.2)                      | 358 (4.6)          | <0.001                                        | 344 (5.1)          | 0.785        |
| Diabetes mellitus               | 1694 (22.5)                    | 1733 (22.2)        | 0.739                                         | 1477 (22.0)        | 0.648        |
| Heart failure history           | 120 (1.6)                      | 71 (0.9)           | <0.001                                        | 65 (1.0)           | 0.795        |
| Renal failure history           | 84 (1.1)                       | 85 (1.1)           | 0.949                                         | 74 (1.1)           | 0.802        |
| Previous MI                     | 575 (7.6)                      | 453 (5.8)          | <0.001                                        | 395 (5.9)          | 0.561        |
| Previous PCI or CABG            | 446 (5.9)                      | 519 (6.7)          | 0.063                                         | 391 (5.8)          | 0.487        |
| Preadmission medication         |                                |                    |                                               |                    |              |
| Aspirin                         | 1288 (17.1)                    | 1259 (16.1)        | 0.126                                         | 1081 (16.1)        | 0.607        |
| P2Y12 receptor inhibitors      | 935 (12.4)                     | 929 (11.9)         | 0.372                                         | 794 (11.9)         | 0.507        |
| Statins                         | 908 (12.0)                     | 815 (10.4)         | 0.002                                         | 714 (10.7)         | 0.523        |

Data are expressed as mean±SD or \( n \) (%). CABG indicates coronary artery bypass grafting; MI, myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST-segment–elevation myocardial infarction.
were found in most subgroups except for MACE risk by the type of MI \((P=0.002\) for interaction) and the disease history of previous percutaneous coronary intervention or coronary artery bypass grafting \((P=0.049\) for interaction); the associated risk of MACE was decreased more significantly after CPC accreditation in patients diagnosed with STEMI \((HR: 0.70; 95\% CI, 0.58–0.84)\) compared with those diagnosed with non-STEMI \((HR: 0.97; 95\% CI, 0.75–1.26)\).

### Table 2. In-Hospital Outcomes Within 7 Days After Hospitalization*

| Variable               | Unmatched Before Accreditation | After Accreditation | Propensity Score Matched Before Accreditation | After Accreditation | \(P\) Value Before Accreditation | \(P\) Value After Accreditation |
|------------------------|-------------------------------|---------------------|----------------------------------------------|---------------------|--------------------------------|--------------------------------|
| n                      | 7544                          | 7800                | 6700                                         | 6700                |                                 |                                |
| MACE, n (%)            | 636 (8.4)                     | 498 (6.4)           | <0.001                                       | 539 (8.0)           | 448 (6.7)                      | 0.003                          |
| All-cause death, n (%) | 120 (1.6)                     | 88 (1.1)            | 0.016                                        | 107 (1.6)           | 75 (1.1)                       | 0.021                          |
| Cardiac death, n (%)   | 114 (1.5)                     | 85 (1.1)            | 0.025                                        | 101 (1.5)           | 72 (1.1)                       | 0.032                          |
| Reinfarction, n (%)    | 18 (0.2)                      | 11 (0.1)            | 0.228                                        | 14 (0.2)            | 10 (0.1)                       | 0.540                          |
| Stent thrombosis, n (%)| 10 (0.1)                      | 7 (0.1)             | 0.579                                        | 9 (0.1)             | 7 (0.1)                        | 0.802                          |
| Stroke, n (%)          | 12 (0.2)                      | 8 (0.1)             | 0.456                                        | 11 (0.2)            | 7 (0.1)                        | 0.479                          |
| Heart failure, n (%)   | 547 (7.3)                     | 419 (5.4)           | <0.001                                       | 459 (6.9)           | 378 (5.6)                      | 0.004                          |

Data are expressed as n (%). MACE indicates major adverse cardiovascular events. *Patients may have had >1 outcome in each category but were counted only once for overall events.

Figure 2. Cumulative Kaplan–Meier curve estimates of outcomes within 7 days after hospitalization. Data for MACE in the whole study population (A) and the propensity score–matched population (B). Data for all-cause death in the whole study population (C) and the propensity score-matched population (D). HR indicates hazard ratio; MACE, major adverse cardiovascular events. DOI: 10.1161/JAHA.119.013384
Sensitivity Analyses

Sensitivity analyses included 36,911 patients admitted to hospitals without accreditations during the project and 8,858 patients contemporarily admitted in hospitals after accreditation (Figure S4). After propensity score matching at a 1:1 ratio, postmatching absolute standardized differences were <10% for all covariates except for admission date. The mirrored histograms are shown in Figure S5 to present the propensity score distribution status.

Baseline characteristics between no accreditation and postaccreditation groups before propensity score matching were not significantly different from the main results except those regarding hospital location, type of MI, and history of diabetes mellitus (Table S1). The results for the risk of in-hospital outcomes within 7 days after admission did not significantly change except for the comparison of all-cause death, with a \( P \) value not significantly different (Tables S2 and S3; Figure S6).

Discussion

Our study found that AMI patients admitted to hospitals with CPC accreditation had better in-hospital outcomes than those admitted to hospitals without accreditation, based on a large-scale national registry data set, even after propensity score matching and using Cox proportional hazards models. To our knowledge, this study is the first to focus on this issue in China. This is especially important because a previous study found no improvement of outcome for STEMI patients from 2001 to 2011. The results of this study confirm the importance and effectiveness of CPC development and accreditation for AMI patients in China.

The most compelling reason for hospitals and programs to obtain accreditations is better patient outcomes, but prior studies focusing on the impact of accreditation were inconclusive. A recent study including >4 million patients aged \( \geq 65 \) years showed no patient mortality benefit for hospitals with a Joint Commission certification compared with those accredited by another independent accrediting organization. However, regarding AMI patients, previous studies have shown that CPC accreditation was associated with better performance on core measurements, which is the fundamental goal for CPC accreditation. These findings suggested that the improvements in care processes with CPC accreditation are further associated with improved short-term outcomes for AMI patients.

Our study found that CPC accreditation was associated with statistically significant risk reduction of MACE and all-

Table 3. Outcomes Within 7 Days After Hospitalization Associated With Accreditation: Unadjusted and Multivariate Adjusted Analyses With and Without Propensity Score Matching

| Variable                        | Unmatched Before Accreditation | Matched Before Accreditation | Propensity Score Matched Before Accreditation | Matched After Accreditation |
|---------------------------------|--------------------------------|------------------------------|-----------------------------------------------|----------------------------|
| MACE                            | Crude HR (95% CI)              | 1.00 (0.79–0.9)              | 1.00 (0.74–0.98)                               | 0.85 (0.78–0.98)           |
| \( P \) value                   | 0.001                          | 0.029                        |                                               |                            |
| Age- and sex-adjusted HR (95% CI)| 1.00 (0.75–0.86)              | 1.00 (0.7–0.94)              |                                               |                            |
| \( P \) value                   | <0.001                         | 0.006                        |                                               |                            |
| Multivariate adjusted HR (95% CI)*| 1.00 (0.77–0.88)              | 1.00 (0.68–0.91)             |                                               |                            |
| \( P \) value                   | <0.001                         | 0.001                        |                                               |                            |

All-cause death

| Variable                        | Unmatched Before Accreditation | Matched Before Accreditation | Propensity Score Matched Before Accreditation | Matched After Accreditation |
|---------------------------------|--------------------------------|------------------------------|-----------------------------------------------|----------------------------|
| Crude HR (95% CI)              | 1.00 (0.67–0.91)              | 1.00 (0.48–0.93)             |                                               |                            |
| \( P \) value                   | 0.011                          | 0.017                        |                                               |                            |
| Age- and sex-adjusted HR (95% CI)| 1.00 (0.64–0.88)              | 1.00 (0.46–0.89)             |                                               |                            |
| \( P \) value                   | 0.005                          | 0.009                        |                                               |                            |
| Multivariate-adjusted HR (95% CI)*| 1.00 (0.69–0.95)              | 1.00 (0.51–0.99)             |                                               |                            |
| \( P \) value                   | 0.022                          | 0.042                        |                                               |                            |

HR indicates hazard ratio; MACE, major adverse cardiovascular events; MI, myocardial infarction.

*Adjusted for age, sex, the level of the city where the hospital is located, first medical contact site or not, comorbidities including smoking status, diabetes mellitus, hypertension, dyslipidemia, diabetes mellitus, heart failure history, renal failure history, previous MI, previous percutaneous coronary intervention or coronary artery bypass grafting, type of MI, Killip classes, preadmission use of aspirin, preadmission use of P2Y12 receptor inhibitors, and preadmission use of statins.

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cause death after adjustment. Although the main contributor to MACE improvement was heart failure, all-cause mortality was also significantly improved with CPC accreditation, even in Cox proportional hazards models, which means the conclusion was solid. Although the history of CPC accreditation in China is relatively short, this study suggested that there were favorable associations with clinical outcomes. Further implementation and optimization of accreditation standards, focused especially on core measurements for care management of AMI patients, may be warranted.

In 2010, the China Expert Consensus on the Construction of Chest Pain Centers was published and played a crucial role in promoting the development of CPCs at that time. The first CPC in China with the regional collaborative network as its core concept was established in March 2011 in Guangzhou. With the scope of collaboration with resources and promoting the rapid development of CPCs, the China Cardiovascular Association and the Chinese Society of Cardiology collaboratively established the China Chest Pain Center in July 2016. After its establishment, this CPC assumed responsibility for most of the organization, management, and coordination originally undertaken by the China Chest Pain Center Accreditation Office. More details on the process, benchmarks, requirements, and so forth, according to CPC certification in China are provided in Data S1 for reference.

An important insight from this study is that the hazards for the risk of outcomes during the in-hospital period as a function of the duration of accreditation followed a reverse J-shaped trend, with the maximum associated effectiveness occurring up until the first year after accreditation. This finding supports the rationale of the recently established reaccreditation project, which may promote a long-lasting effect after initial CPC accreditation and help to limit subsequent declines in associated benefits for AMI care and outcomes with CPC accreditation after the first year. In January 2018, considering continuing quality improvement, the headquarters of China CPCs formally established a reaccreditation protocol and standard that demands a 3-year validation period for the initial CPC accreditation. The validation period for the reaccreditation of CPCs will be 5 years, and reaccreditation should be performed every 5 years.

This study had several limitations. First, this was a retrospective observational study; even using propensity score matching, we could not eliminate bias from unobserved

Figure 3. Hazard for the risk of outcomes within 7 days after hospitalization by the duration of accreditation. Data for MACE in the whole study population (A) and the propensity score–matched population (B). Data for all-cause death in the whole study population (C) and the propensity score–matched population (D). MACE indicates major adverse cardiovascular events.
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variables. A follow-up study with a more robust design, such as prospective cluster randomization, would be warranted. Second, the difference in outcomes was largely driven by difference in heart failure and all-cause death, which should be taken into consideration in the interpretation of results. Third, hospitals involved in CCC-ACS projects cannot reflect the whole picture of China’s hospitals with and without CPC accreditation; furthermore, this study focuses on only AMI rather than all acute chest pain diagnoses and diseases. Last, our analysis was able to focus on only in-hospital outcomes rather than effect of accreditation on long-term outcome improvements, which can be further studied in the future.

Conclusions

Based on a large-scale national registry data set, CPC accreditation was associated with better in-hospital outcomes for AMI patients. However, because there was some attenuation in associated benefits over time, reaccreditation may be essential to maintain the quality of AMI care after initial CPC accreditation.

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Disclosures

None.

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SUPPLEMENTAL MATERIAL
Appendix S1 Group information: *Hospitals for Phase One (Investigator)*:

Shanxi Cardiovascular Hospital (Bao Li); Nanjing Drum Tower Hospital, The Affiliated Hospital of Nanjing University Medical School (Biao Xu, Guangshu Han); Hainan General Hospital (Bin Li); The Second Hospital of Jilin University (Bin Liu); The 2nd Affiliated Hospital of Harbin Medical University (Bo Yu); The Ninth Hospital Affiliated to Shanghai Jiaotong University School of Medicine (Changqian Wang); Henan Provincial People's Hospital (Chuanyu Gao); Shanxi Provincial People's Hospital (Chunlin Lai); Xinqiao Hospital, Third Military Medical University (Cui Bin, Lan Huang); China Meitan General Hospital (Di Wu); The 309th Hospital of Chinese People's Liberation Army (Fakuan Tang, Jun Xiao); Zhongda Hospital, Southeast University (Genshan Ma); The First Affiliated Hospital of Liaoning Medical University (Guizhou Tao); Xinjiang Uygur Autonomous Region People’s Hospital (Guoqing Li); Sir Run Run Shaw Hospital, College of Medicine, Zhejiang University (Guosheng Fu); Beijing Friendship Hospital, Capital Medical University (Hongwei Li); The First Affiliated Hospital of Bengbu Medical College (Honhju Wang); General Hospital of TISCO (Huifeng Wang); Dongguan People's Hospital (Jianfeng Ye); Panyu Hospital of Chinese Medicine (Jianhao Li); Peking University First Hospital (Jie Jiang); Sun Yat-sen Memorial Hospital, Sun Yat-sen University (Jingfeng Wang); Guangdong General Hospital (Jiyan Chen); Hospital of Xinjiang Production & Construction Corps (Junming Liu); The Military General Hospital of Beijing PLA (Junxia Li); The First Affiliated Hospital of Guangxi Medical University (Lang Li); Tongren Hospital Affiliated to Shanghai Jiaotong University School of Medicine (Li Jiang); Binzhou City Center Hospital (Lijun Meng); The First Affiliated Hospital of Zhengzhou University (Ling Li); Xijing
Hospital (Ling Tao); The Affiliated Hospital of Guizhou Medical University (Lirong Wu); First Affiliated Hospital of the People's Liberation Army General Hospital (Miao Tian); The Second People's Hospital of Yunnan Province (Minghua Han); Haikou People's Hospital (Moshui Chen); Gansu Provincial Hospital (Ping Xie); The First Affiliated Hospital of Henan University of Science and Technology (Pingshuan Dong); Chenzhou First People's Hospital (Qiaoqing Zhong); People’s Hospital of Qinghai Province (Rong Chang); Affiliated Hospital of Ningxia Medical University (Shaobin Jia); Beijing Anzhen Hospital, Capital Medical University (Shaoping Nie, Xiaohui Liu); North Jiangsu People's Hospital (Shenghu He); Shanghai Sixth People's Hospital (Shixin Ma); The First Hospital of Handan (Shuanli Xin); Huai'an First People's Hospital (Shuren Ma); The First Affiliated Hospital of Chongqing Medical University (Suxin Luo); Navy General Hospital (Tianchang Li); Zhejiang Provincial Hospital of TCM (Wei Mao); The Third Xiangya Hospital of Central South University (Weihong Jiang); Affiliated Hospital of Qinghai University (Weijun Liu); Teda International Cardiovascular Hospital (Wenhua Lin); The Second Hospital of Hebei Medical University (Xianghua Fu); Changhai Hospital of Shanghai (Xianxian Zhao); The Second Affiliated Hospital to Nanchang University (Xiaoshu Cheng); Hebei General Hospital (Xiaoyong Qi); Inner Mongolia People's Hospital (Xingsheng Zhao); The General Hospital of Shenyang Military Region (Yaling Han); The First Hospital of Jilin University (Yang Zheng); Tianjin Chest Hospital (Yin Liu); Hunan Provincial People's Hospital (Ying Guo); People's Hospital of Yuxi City (Yinglu Hao); The People's Hospital of Guangxi Zhuang Autonomous Region (Yingzhong Lin); The First Teaching Hospital of Xinjiang Medical University (Yitong Ma); Baogang Hospital (Yongdong Li); Tianjin Medical University General Hospital (Yuemin
Sun); The Second Affiliated Hospital of Zhengzhou University (Yulan Zhao); Nanfang Hospital of Southern Medical University (Yuqing Hou); The First Affiliated Hospital to Nanchang University (Zeqi Zheng); The First Affiliated Hospital of Lanzhou University (Zheng Zhang); The Third Hospital of Shijiazhuang (Zhenguo Ji); Wuxi People's Hospital (Zhenyu Yang); Jiangsu Province Hospital (Zhijian Yang); The Second Hospital of Shanxi Medical University (Zhiming Yang); The Affiliated Hospital of Xuzhou Medical College (Zhirong Wang); Southwest Hospital, Third Military Medical University (Zhiyuan Song); The First Affiliated Hospital of Xi’an Jiaotong University (Zuyi Yuan). \textit{Hospitals for Phase Two (Investigator)}: Yangzhou First People's Hospital (Aihua Li); Hospital 463 of Chinese People's Liberation Army (Bosong Yang); The Central Hospital of Mianyang (Caidong Luo); Liaocheng People's Hospital (Chunyan Zhang); Yancheng Third People's Hospital (Chunyang Wu); The Second Xiangya Hospital of Central South University (Daoquan Peng); The Central Hospital of Panzhihua (Dawen Xu); The First Hospital of Qiqihaer City (Gang Xu); The Third the People’s Hospital of Bengbu (Gengsheng Sang); The First Hospital of Jiamusi (Guixia Zhang); Zhoushan People's Hospital (Guoxiong Chen); Dalian Municipal Central Hospital (Hailong Lin); Renmin Hospital of Wuhan University (Hong Jiang); Ningxia People's Hospital (Hong Luan); The First People's Hospital of Yunnan Province (Kunhua Hospital) (Hong Zhang); The Central Hospital of Zhoukou (Hualing Liu); Anyang District Hospital (Hui Liu); Sichuan Provincial People’s Hospital (Jianhong Tao); Mudanjiang Cardiovascular Disease Hospital (Jianwen Liu); Yichang Central Hospital (Jiawang Ding); Qilu Hospital of Shandong University (Jifu Li); Affiliated Hospital of Jiangsu University (Jinchuan Yan); The First People's Hospital of Nanning City (Jinru Wei); The First Affiliated
Affiliated Hospital of Wannan Medical College (Xingsheng Tang); Tangdu Hospital of The Fourth Military Medical University (Xue Li); Shanghai East Hospital Affiliated to Tongji University (Xuebo Liu); Xiamen Cardiovascular Disease Hospital (Yan Wang); Zhongnan hospital of Wuhan University (Yanggan Wang); Fujian Provincial Hospital (Yansong Guo); The First Affiliated hospital of Dalian Medical University (Yanzong Yang); The First People's Hospital of Changde (Yi Huang); The First Affiliated Hospital of China Medical University (Yingxian Sun); The Fourth Affiliated Hospital of China Medical University (Yuanzhe Jin); Cangzhou Central Hospital (Zesheng Xu); The Central Hospital of Shaoyang (Zewei Ouyang); The People's Hospital of Liaoning Province (Zhanquan Li); The First Affiliated Hospital of Jiamusi University (Zhaofa He); Tangshan Gongren Hospital (Zheng Ji); Huaibei Miners General Hospital (Zhenqi Su); Linyi People's Hospital (Zhihong Ou).
Data S1. Details on the process, benchmarks, requirements according to the CPC accreditation in China

The accreditation standard system of CPC in China

1. CPC Accreditation Standard: China Chest Pain Center Accreditation Standard was released in 2013, including a standard version and a basic version for different hospitals. The standard version is suitable for the annual pPCI operation volume ≥200 cases and guarantee 7x24 hours for emergency PCI operation. The main target of the basic version is for those hospitals do not meet the requirements of the standard version, but the number of patients with acute myocardial infarction who are admitted/referred to ≥ 30 cases per year. The two sets of standards effectively cover the needs of the construction of CPCs in China. With the continuous accumulation of accreditation work experience, the actual situation of medical institutions in different regions of the country has been comprehensively revised. The standard has been revised several times during the period. The current standard version has been updated to the fifth edition, and the basic version has been updated to the second in version.

2. Re-accreditation standard: In order to continuously supervise and manage the certified units, the accreditation standard clearly requires that the CPC should be valid for 3 years after the first accreditation, and each CPC needs to be re-accredited before the expiration date. In January 2018, the China Chest Pain Center Accreditation Working Committee issued the “China Chest Pain Center Re-accreditation Standard (Standard Edition)”. With the approval of the first batch of primary CPCs at the end of 2016, the re-accreditation standards for basic CPCs will be drafted and announced by the end of 2019.

3. Construction standards: In order to further clarify the basic requirements for the construction of a CPC, assist the hospital to conduct self-assessment, and encourage more medical institutions to join the CPC construction team. In September 2016, the China Chest Pain Center Accreditation Work Committee drafted, revised and released "standards for the construction of chest pain centers”.

4. Group standards: In order to actively respond to the call for reforming of the national
standardization system, effectively promote the implementation of the policies of the National Health and Health Commission, and provide a solid foundation for the future CPC network in China, in August 2018, "China Cardiopulmonary Resuscitation and Cardiac Defibrillation Group Standards and China Chest Pain Center Construction and Evaluation Group Standards", which was prepared by the Rescue Association, the Chinese Nursing Association, the China Cardiovascular Health Alliance, and the Cardiovascular Health (Suzhou Industrial Park) Institute, was officially released and published on the national group standard information platform (standard numbers T/CADERM2001-2018 and T/CADERM2002-2018).

**Construction and review process of CPC in China**

Medical institutions at all levels actively responded to the policy requirements of the government, promote the construction of CPCs, and provide patients with higher quality medical services. They need to go through three stages before the accreditation, namely self-construction stage, the review phase and the continuous quality improvement phase.

1. **Self-construction:** The establishment of a CPC committee should led by the dean or the deputy medical dean of the medical institution, covering all the heads of medical and technical departments related to acute chest pain treatment, and also setting up medical directors, administrative directors, and liaisons. The person in charge of the position, conduct self-assessment, and in accordance with the requirements of the published accreditation standards (standard or basic version), by participating in relevant construction training sessions or watching the operation experience of the demonstration base, set up corresponding points in the main roads and major treatment points. The identification/guidance, optimize the relevant treatment process, unify the hospital clock, sign a joint treatment agreement with the pre-hospital emergency and network hospital, classify the whole staff treatment training in the hospital and the community, and faithfully record all except traumatic chest pain.

2. **Review phase:** Data on treatment of patients with acute chest pain and training related raw materials, and continuous optimization of the treatment process based on phased data analysis, when the CPCs actually running for at least 6 months (providing no less than 6 months of data
to assess the quality of operation) and self-assessment after basically meeting the requirements of the accreditation standard, at the specified time in the inter-node, the application for accreditation is issued to the China Chest Pain Center Accreditation Working Committee and the Chest Pain Center Headquarters. The whole year is divided into 4 batches of accreditation stages. After receiving the accreditation application from the medical institution, the headquarters first checks the accreditation qualification (including the consultation). Whether the quantity meets the requirements of the corresponding standard, whether the construction materials are complete, whether the case record meets the requirements of 6 months, whether the key treatment indicators are up to standard, etc.), the unit that passed the examination is handed over to the corresponding regional accreditation office for detailed review of each accreditation clause, and the unsatisfied clause proposed rectification opinions. After the unit was rectified, the office re-evaluated and recommended to enter the provincial pre-examination list of the provincial chest pain center. The provincial alliance assigned the provincial pre-inspection experts to the hospital for field visits, and conducted the operation of the chest pain center of the applicant unit. After the actual evaluation and suggestions for improvement, the provincial alliance comprehensively gives the results of the batch pre-inspection after passing the pre-inspection or rectification. Units that pass the pre-inspection of the provincial alliance will accept random and anonymous online materials and data reviews from three non-provincial national verification experts. Applicants who pass the expert online review will enter the unannounced visit and on-site verification stage. The unannounced visit is an expert commissioned by the headquarters to investigate the hospital's identification guidelines without the knowledge of the accreditation unit, and observe the applicant's actual acute chest pain patients or hire volunteers to simulate the patient's diagnosis and treatment. The diagnosis process and assessment of whether each link is standardized, and the corresponding score. The on-site inspection was carried out by three non-provincial experts and an office staff member to the applicant to evaluate the construction materials of the chest pain center, the authenticity of the data and the training effect of the whole staff, and to give a rating. All the online evaluations, unannounced visits and on-site inspections and scores of the applicants entering
the unannounced visit/site verification stage will be reported by the office at the plenary session of the China Chest Pain Center Accreditation Working Committee and will be voted by secret ballot on the overall situation of the members present. More than half of the votes will be passed, otherwise they will need to be revisited and/or checked again. The units that have passed the voting will elect to issue the bronze medals that have passed the accreditation.

3. **Continuous quality improvement**: Continuous improvement is the standard operation of the CPC, ensuring the essence of providing high-efficiency treatment. For certified CPCs, accreditation standards should continue to be optimized, enhanced training and extensive publicity and education, and accurate data management to further improve the efficiency and level of treatment for acute chest pain-related diseases. The China Chest Pain Center Accreditation Working Committee has separately formulated the "China Chest Pain Center Quality Control Indicators and Assessment Methods" (Standard Edition and Basic Edition, now updated to the second edition) and "China Chest Pain Center Re-accreditation Standards" (first edition) promote continuous improvement of accreditation through the unit. Specifically, the quality control center will monitor and evaluate the operation quality of each chest pain center every month. The regional reaccreditation office will conduct an overall assessment of the operation of the chest pain center within three years after accreditation.

**Quality control system of China CPC**

"Without scale, there is no benefit, no quality, no life". In keeping with this aim, while promoting the rapid growth of the number of CPCs, the Accreditation Work Committee and the Chest Pain Center Headquarter continue to promote and improve quality control methods.

1. **Establish a data platform for chest pain center data and provide it to all chest pain centers for free.** The patient type collected by the data reporting platform covers all patients with acute chest pain except traumatic chest pain. The collected fields focus on the treatment time node and key medical behavior record, providing data support for accreditation, quality control and reaccreditation assessment. Current data reporting platform In the process of upgrading, it is mainly updated from two angles. The first is the classification of data fields.
The existing fields are summarized into the basic information of the patient, the basic information of the disease and the information of the specialist diagnosis and treatment, for the future and the center of atrial fibrillation and heart failure. The integration of data platforms such as rehabilitation centers has paved the way for the formation of a big data center for cardiovascular diseases, which has become an important support for government decision-making and the development of the industry. Second, the positioning of database roles, providing basic library of accreditation quality control and scientific research to meet different positioning. The work needs of the medical institutions have made the chest pain center database truly an assistant to departmental patient management and to promote medical quality improvement.

2. Promote the development of data collection information, improve reporting efficiency and data quality. Develop existing real-time filing and data uploading applets, apps or automatic collection technology with existing Internet technologies, to reduce the workload of data manual reporting, reduce error rate, improve data quality and report efficiency, and provide reliable quality control.

3. Establish a data quality control platform for the chest pain center to provide a grasp for the provincial/local alliance to promote quality control. The establishment of the quality control platform will summarize and analyze the data reported by the CPCs of each unit, and present them in a visualized form to express the continuous quality improvement of each CPC. The index ranking has become an important starting point for the provincial/municipal chest pain center alliance to carry out quality control work.

4. Establish a normalized quality control system to promote the continuous improvement of the quality of the chest pain center. The normalized quality control system includes: issuing quality control indicators and assessment methods for chest pain centers, developing quality control platforms, formulating quality control reports for health care committees, provincial/municipal chest pain center alliances, hospitals, etc.. The system is commended and criticized at the annual Chest Pain Center Quality Control Conference. Thereby establishing a four-level quality control system for hospital internal quality control, municipal-level alliance
quality control, provincial-level alliance quality control and national quality control. In the future, with the support of the National Health and Health Commission, the quality control effect of the chest pain center can provide data support for the grade hospitals or key disciplines, and can also provide reference for the cost settlement of the medical insurance bureaus at all levels.
Table S1. Baseline Characteristics of the Study Population in Sensitivity Analyses.

| Variables                        | Unmatched |                  |                | Propensity score-matched |                  |                |
|----------------------------------|-----------|------------------|----------------|--------------------------|------------------|----------------|
|                                  | no        | after            | p value        | no                       | after            | p value        |
|                                  | accreditation | accreditation |                | accreditation            | accreditation |                |
| n                                | 36911     | 8858             | 0.21           | 8719                     | 8719            | 0.59           |
| Age, years                       | 62.66 (12.66) | 62.48 (12.44)     | 1              | 62.36 (12.50)            | 62.46 (12.45)   | 0.34           |
| Sex: Female                      | 8862 (24.0) | 2100 (23.7)       | 0.55           | 2015                     | 2069 (23.7)     | 3              |
| Hospital location                |           |                  |                |                          |                  |                |
| First-line municipality          | 5715 (15.5) | 0 (0.0)         |                |                          |                  |                |
| Provincial capital               | 15297 (41.4) | 2798 (31.6)       | 0.30           | 5985                     | 5921 (67.9)     |                |
| Prefecture-level city            | 15899 (43.1) | 6060 (68.4)       |                | 2734                     | 2798 (32.1)     |                |
| Hospital: First medical facility | 19223 (52.1) | 4844 (54.7)       | <0.00          | 4790                     | 4745 (54.4)     | 0.50           |
| Department: Emergency/Cath lab    | 24039 (65.1) | 6797 (76.7)       | <0.00          | 6507                     | 6660 (76.4)     | 0.00           |
| Type of MI                       |            |                  |                |                          |                  |                |
| STEMI                            | 26620 (72.1) | 6303 (71.2)       | 0.07           | 6104                     | 6207 (71.2)     | 0.09           |
| Killip class                     |            |                  | <0.00          |                          |                  | 0.97           |
| Class I                          | 25828 (70.0) | 6584 (74.3)       |                | 6444                     | 6457 (74.1)     |                |
| Class II-III                     | 9394 (25.5) | 1970 (22.2)       |                | 1971                     | 1959 (22.5)     |                |
| Class IV                         | 1689 (4.6) | 304 (3.4)         |                | 304 (3.5)                | 303 (3.5)       |                |
| Comorbidity                      |            |                  |                |                          |                  |                |
| Current smoking                  | 16183 (43.8) | 4193 (47.3)       | <0.00          | 4099                     | 4103 (47.1)     | 0.96           |
| Hypertension                     | 19013 (51.5) | 4495 (50.7)       | 0.71           | 4461                     | 4422 (50.7)     | 0.56           |
| Dyslipidemia                     | 2922 (7.9) | 427 (4.8)         | 0.20           | 463 (5.3)                | 426 (4.9)       | 0.21           |
| Diabetes mellitus                | 7999 (21.7) | 1936 (21.9)       | <0.00          | 1880                     | 1903 (21.8)     | 0.68           |
| Condition                        | Mean ± SD | n (%) | p-value |
|---------------------------------|-----------|-------|---------|
| Heart failure history           | 762 (2.1) | 85 (1.0) | <0.01 |
| Renal failure history           | 701 (1.9) | 93 (1.0) | <0.01 |
| Previous MI                     | 2650 (7.2)| 512 (5.8)| <0.01 |
| Previous PCI or CABG            | 2298 (6.2)| 584 (6.6)| 0.21  |
| **Pre-admission of medication** |           |       |         |
| Aspirin                         | 8214 (22.3)| 1467 (16.6) | <0.01 |
| P2Y<sub>12</sub> receptor inhibitors | 6176 (16.7)| 1105 (12.5) | <0.01 |
| Statins                         | 5737 (15.5)| 1008 (11.4) | <0.01 |

Data are expressed as means ± SD, or n (%). Abbreviations: CABG, coronary artery bypass graft; Cath, Catheter; MI, myocardial infarction; NSTEMI, non-ST elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST elevation myocardial infarction.
Table S2. In-hospital Outcomes* within 7 days after Hospitalization in Sensitivity Analyses.

| Variables                | Unmatched                               | Propensity score-matched      | p value | Unmatched                               | Propensity score-matched      | p value |
|--------------------------|-----------------------------------------|--------------------------------|---------|-----------------------------------------|--------------------------------|---------|
|                          | no accreditation | after accreditation | p value | no accreditation | after accreditation | p value |
| n                        | 36911                      | 8858                        | <0.001  | 8719                      | 8719                        | <0.001  |
| MACE, n(%)               | 4376 (11.9)                | 542 (6.1)                   | <0.001  | 810 (9.3)                | 534 (6.1)                   | <0.001  |
| All-cause death, n(%)    | 621 (1.7)                  | 107 (1.2)                   | 0.002   | 121 (1.4)                | 106 (1.2)                   | 0.350   |
| Cardiac death, n(%)      | 591 (1.6)                  | 104 (1.2)                   | 0.004   | 114 (1.3)                | 103 (1.2)                   | 0.495   |
| Re-infarction, n(%)      | 88 (0.2)                   | 15 (0.2)                    | 0.268   | 24 (0.3)                 | 15 (0.2)                    | 0.200   |
| Stent thrombosis, n(%)   | 50 (0.1)                   | 8 (0.1)                     | 0.365   | 7 (0.1)                  | 8 (0.1)                     | 1.000   |
| Stroke, n(%)             | 112 (0.3)                  | 9 (0.1)                     | 0.001   | 26 (0.3)                 | 9 (0.1)                     | 0.007   |
| Heart failure, n(%)      | 3907 (10.6)                | 440 (5.0)                   | <0.001  | 696 (8.0)                | 433 (5.0)                   | <0.001  |

Data are expressed as n (%). Abbreviations: MACE, major adverse cardiovascular events.

*Patients may have had more than 1 outcome in each category but counted only once for overall events.
Table S3. Independent Predictors of MACE in Propensity Score-matched Population in Sensitivity Analyses.

| Variables                        | Unmatched | Propensity score-matched |
|----------------------------------|-----------|--------------------------|
|                                  | Before accreditation | After accreditation | Before accreditation | After accreditation |
| MACE                             |            |                          |                      |
| Crude HR (95% CI)                | 1.00       | 0.50 (0.46, 0.55)        | 1.00                 | 0.65 (0.58, 0.73)   |
| p value                          | <0.001     |                          | <0.001               |                      |
| Age and sex adjusted HR (95% CI)| 1.00       | 0.50 (0.46, 0.55)        | 1.00                 | 0.64 (0.58, 0.72)   |
| p value                          | <0.001     |                          | <0.001               |                      |
| Multivariate adjusted HR         | 1.00       | 0.57 (0.52, 0.62)        | 1.00                 | 0.64 (0.58, 0.72)   |
| p value                          | <0.001     |                          | <0.001               |                      |
| All-cause death                  |            |                          |                      |
| Crude HR (95% CI)                | 1.00       | 0.72 (0.59, 0.89)        | 1.00                 | 0.88 (0.68, 1.15)   |
| p value                          | 0.002      |                          | 0.350                |                      |
| Age and sex adjusted HR (95% CI)| 1.00       | 0.73 (0.60, 0.90)        | 1.00                 | 0.87 (0.67, 1.14)   |
| p value                          | 0.003      |                          | 0.314                |                      |
| Multivariate adjusted HR (95% CI)| 1.00      | 0.73 (0.59, 0.90)        | 1.00                 | 0.87 (0.67, 1.13)   |
| p value                          | 0.003      |                          | 0.289                |                      |

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; HR, hazard ratio; MACE, major adverse cardiovascular events; MI, myocardial infarction; PCI, percutaneous coronary intervention.

* Adjusted for age, sex, the level of the city where the hospital is located, first medical contact site or not, comorbidities including smoking status, diabetes mellitus, hypertension, dyslipidemia, diabetes mellitus, heart failure history, renal failure history, previous MI, previous PCI or CABG, type of MI, Killip classes, pre-admission use of aspirin, pre-admission use of P2Y12 receptor inhibitors, and pre-admission use of statins.
Figure S1. Mirrored Histogram before (A) and after (B) propensity score matching. X axis is the number of patients in each group. Y axis is the propensity score. The red bar presents the before accreditation group and the blue bar for the after accreditation group.
Figure S2. Subgroup analyses for MACE in the propensity score–matched population. Abbreviations: CABG, coronary artery bypass graft; Cath, Catheter; CI, confidence interval; MACE, major adverse cardiovascular events; MI, myocardial infarction; NSTEMI, non-ST elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST elevation myocardial infarction.

| Subgroups          | No. of Patients | Before | After | Hazard Ratio (95%CI) | p value | p for interaction |
|--------------------|-----------------|--------|-------|----------------------|---------|------------------|
| Overall            | 13400           | 539 (8.0) | 448 (6.7) |                      | 0.78 (0.68, 0.81) | 0.001            |
| Age                |                 |        |       |                      |         |                  |
| <65 year           | 7457            | 177 (4.9) | 152 (4.1) | 0.80 (0.61, 1.04)   | 0.980   | 0.780            |
| ≥65 year           | 5943            | 362 (12.2) | 298 (10.0) | 0.79 (0.67, 0.95)   | 0.036   |                  |
| Sex                |                 |        |       |                      |         |                  |
| Male               | 10270           | 342 (6.7) | 298 (5.8) | 0.80 (0.68, 0.95)   | 0.089   | 0.780            |
| Female             | 3130            | 197 (12.6) | 150 (9.6) | 0.76 (0.59, 0.97)   | 0.250   |                  |
| City               |                 |        |       |                      |         |                  |
| Prefecture-level city | 4859      | 274 (11.1) | 179 (7.5) | 0.69 (0.55, 0.88)   | 0.037   | 0.200            |
| Provincial capital | 8541            | 265 (6.2) | 269 (6.3) | 0.85 (0.70, 1.03)   | 0.150   |                  |
| First medical contact site |   |        |       |                      |         |                  |
| No                 | 6667            | 276 (8.3) | 227 (6.8) | 0.74 (0.60, 0.91)   | 0.100   | 0.110            |
| Yes                | 6733            | 263 (7.8) | 221 (6.6) | 0.84 (0.68, 1.03)   | 0.220   |                  |
| Department arrived |                 |        |       |                      |         |                  |
| Outpatient, others | 3380            | 141 (8.4) | 101 (5.9) | 0.79 (0.57, 1.09)   | 0.160   | 0.850            |
| Emergency, cath lab| 10020           | 398 (7.9) | 347 (6.9) | 0.80 (0.67, 0.94)   | 0.110   |                  |
| Type of MI         |                 |        |       |                      |         |                  |
| STEMI              | 3745            | 166 (8.8) | 160 (8.6) | 0.70 (0.58, 0.84)   | 0.140   | 0.002            |
| NSTEMI             | 9655            | 373 (7.7) | 288 (6.0) | 0.97 (0.75, 1.26)   | 0.064   |                  |
| Killip class I     |                 |        |       |                      |         |                  |
| No                 | 9714            | 239 (4.9) | 194 (4.0) | 0.74 (0.60, 0.92)   | 0.500   | 0.850            |
| Yes                | 3686            | 300 (16.1) | 254 (13.9) | 0.80 (0.65, 0.97)   | 0.009   |                  |
| Previous MI        |                 |        |       |                      |         |                  |
| No                 | 12593           | 496 (7.9) | 402 (6.4) | 0.77 (0.67, 0.9)    | 0.053   | 0.560            |
| Yes                | 807             | 43 (10.9) | 46 (11.2) | 1.07 (0.63, 1.84)   | 0.830   |                  |
| Previous PCI or CABG |               |        |       |                      |         |                  |
| No                 | 12597           | 509 (8.1) | 407 (6.5) | 0.76 (0.65, 0.88)   | 0.018   | 0.049            |
| Yes                | 803             | 30 (7.7) | 41 (10.0) | 1.16 (0.68, 2.00)   | 0.390   |                  |
Figure S3. Subgroup analyses for all-cause death in the propensity score-matched population. Abbreviations: CABG, coronary artery bypass graft; Cath, Catheter; CI, confidence interval; MACE, major adverse cardiovascular events; MI, myocardial infarction; NSTEMI, non-ST elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST elevation myocardial infarction.

| Subgroups                  | No. of Patients | Before | After | Hazard Ratio (95%CI) p value p for interaction |
|----------------------------|----------------|--------|-------|-----------------------------------------------|
| Overall                    | 13400          | 107 (1.6) | 75 (1.1) | 0.71 (0.51, 0.99) 0.042 0.042 |
| Age                        |                |        |       |                                               |
| <65 year                   | 7457           | 21 (0.6) | 19 (0.5) | 0.99 (0.49, 2.00) 0.980 0.560 |
| ≥65 year                   | 5943           | 88 (2.9) | 56 (1.9) | 0.67 (0.46, 0.97) 0.036 0.036 |
| Sex                        |                |        |       |                                               |
| Male                       | 10270          | 67 (1.3) | 49 (1.0) | 0.68 (0.45, 1.03) 0.069 0.970 |
| Female                     | 3130           | 40 (2.6) | 26 (1.7) | 0.74 (0.44, 1.24) 0.250 0.250 |
| City                       |                |        |       |                                               |
| Prefecture-level city      | 4859           | 47 (1.9) | 23 (1.0) | 0.55 (0.31, 0.96) 0.037 0.510 |
| Provincial capital         | 8541           | 60 (1.4) | 52 (1.2) | 0.74 (0.49, 1.12) 0.150 0.150 |
| First medical contact site |                |        |       |                                               |
| No                         | 6667           | 43 (1.3) | 28 (0.8) | 0.65 (0.39, 1.09) 0.100 0.480 |
| Yes                        | 6733           | 64 (1.9) | 47 (1.4) | 0.77 (0.50, 1.17) 0.220 0.220 |
| Department arrived         |                |        |       |                                               |
| Outpatient, others         | 3380           | 16 (1.0) | 9 (0.5) | 0.48 (0.17, 1.33) 0.160 0.440 |
| Emergency, cath lab        | 10020          | 91 (1.8) | 66 (1.3) | 0.75 (0.52, 1.07) 0.110 0.110 |
| Type of MI                 |                |        |       |                                               |
| STEMI                      | 3745           | 34 (1.8) | 16 (0.9) | 0.75 (0.51, 1.10) 0.140 0.650 |
| NSTEMI                     | 9655           | 73 (1.5) | 59 (1.2) | 0.56 (0.31, 1.03) 0.064 0.064 |
| Killip class I             |                |        |       |                                               |
| No                         | 9714           | 36 (0.7) | 31 (0.6) | 0.84 (0.49, 1.42) 0.500 0.880 |
| Yes                        | 3686           | 71 (3.8) | 44 (2.4) | 0.57 (0.37, 0.87) 0.009 0.009 |
| Previous MI                |                |        |       |                                               |
| No                         | 12593          | 100 (1.6) | 70 (1.1) | 0.71 (0.50, 1.00) 0.053 0.940 |
| Yes                        | 807            | 7 (1.8) | 5 (1.2) | 0.86 (0.23, 3.26) 0.630 0.630 |
| Previous PCI or CABG       |                |        |       |                                               |
| No                         | 12597          | 105 (1.7) | 69 (1.1) | 0.66 (0.47, 0.93) 0.018 0.170 |
| Yes                        | 803            | 2 (0.5) | 6 (1.5) | 2.30 (0.35, 15.03) 0.390 0.390 |
Figure S4. Flow Diagram of Selection of the Study Population in Sensitivity Analyses. Abbreviations: CCC-ACS: Improving Care for Cardiovascular Disease in China-Acute Coronary Syndrome; UAP: Unstable Angina Pectoris.
Figure S5. Mirrored Histogram before (A) and after (B) propensity score matching in Sensitivity Analyses. X axis is the number of patients in each group. Y axis is the propensity score. The red bar presents the no accreditation group and the blue bar for the after accreditation group.
**Figure S6. Cumulative Kaplan-Meier Curve Estimates of Outcomes within 7 days after Hospitalization in Sensitivity Analyses.** Abbreviations: CI, confidence interval; HR, hazard ratio; MACE, major adverse cardiovascular events. Panel A and B show data for MACE in the whole study population and the propensity score-matched population respectively. Panel C and D show data for all-cause death in the whole study population and the propensity score-matched population respectively.