Impact of team-based community healthcare on preventable hospitalisation: a population-based cohort study in Taiwan

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ABSTRACT

Objectives The objective of this study was to explore the impact of Taiwan’s Family Practice Integrated Care Project (FPICP) on hospitalisation.

Design A population-based cohort study compared the hospitalisation rates for ambulatory care sensitive conditions (ACSCs) among FPICP participating and non-participating patients during 2011–2015.

Setting The study accessed the FPICP reimbursement database of Taiwan’s National Health Insurance (NHI) administration containing all NHI administration-selected patients for FPICP enrolment.

Participants The NHI administration-selected candidates from 2011 to 2015 became FPICP participants if their primary care physicians joined the project, otherwise they became non-participants.

Interventions The intervention of interest was enrolment in the FPICP or not. The follow-up time interval for calculating the rate of hospitalisation was the year in which the patient was selected for FPICP enrolment or not.

Primary outcome measures The study’s primary outcome measures were hospitalisation rates for ACSC, including asthma/chronic obstructive pulmonary disease (COPD), diabetes or its complications and heart failure. Logistic regression was used to calculate the ORs concerning the influence of FPICP participation on the rate of hospitalisation for ACSC.

Results The enrolled population for data analysis was between 3.94 and 5.34 million from 2011 to 2015. Compared to non-participants, FPICP participants had lower hospitalisation for COPD/asthma (28.6‰–35.9‰ vs 37.9‰–42.3‰) and for diabetes or its complications (10.8‰–14.9‰ vs 12.7‰–18.1‰) but not for congestive heart failure. After adjusting for age, sex and level of comorbidities by logistic regression, participation in the FPICP was associated with lower hospitalisation for COPD/asthma (OR 0.91, 95% CI 0.87 to 0.94 in 2015) and for diabetes or its complications (OR 0.87, 95% CI 0.83 to 0.92 in 2015).

Conclusion Participation in the FPICP is an independent protective factor for preventable ACSC hospitalisation. Team-based community healthcare programs such as the FPICP can strengthen primary healthcare capacity.

INTRODUCTION

Taiwan’s National Health Insurance (NHI) programme is renowned for its cost-effectiveness and accessibility and serves 23.8 million people with a 99.6% coverage as a high-performing single-payer health insurance system.1,2 Nevertheless, Taiwan also has to face a serious challenge to its financial sustainability due to an ageing population, an insufficient insurance premium rate, as well as fragmented and less patient-centred integrated care as a result of fee-for-service-based payments. To maintain quality care and reduce wasting of resources, Taiwan’s government has been taking action with interventions and policies aimed to reinforce the healthcare capacity of primary care physicians and to re-emphasise general medical training. One major intervention is the establishment of the Family Practice Integrated Care Project (FPICP).3

In brief, the FPICP is a modified pay-for-performance (P4P) programme that affects...
10% of all NHI beneficiaries. Featuring team-based care provided by primary care clinics with integration with community hospitals, the FPICP was started as a pilot project in 2003 and was reformed in 2010 as a regular government healthcare programme. The community healthcare group (CHCG), a team of 5–10 primary care physicians in a single community working in cooperation with a local hospital, forms the core healthcare unit of the FPICP. The target population of the FPICP are patients with multiple chronic diseases, frequent users of outpatient care and the elderly aged over 65 years. Taiwan’s NHI administration selects those incurring higher medical costs among target patients on a yearly basis to compile a list of FPICP candidates. There is no ‘cherry-picking phenomenon’ because the FPICP requires the member physicians to include all administration-assigned patients of the NHI. Primary care physicians of the FPICP deliver integrated healthcare services through collaboration within the CHCG, focusing especially on preventive care, providing continuous care with 24-hour telephone hotline consultation with hospital doctors and a bidirectional mutual referral network among the primary care clinics and local hospitals. If an FPICP participant is hospitalised, the primary care physician can visit the patient in the hospital and participate in the ward round, facilitating further referral back to the primary care clinic. Until the end of 2015, approximately 25% of primary care physicians and 30% of community clinics joined the FPICP to serve the participating patients in 426 CHCGs. Integrated healthcare services provided by primary care physicians of the FPICP are incentivised in addition to a regular fee-for-service payment scheme. On average, each physician is responsible for 550–750 participating FPICP patients. The programme allocates 250 points to member physicians (1 point=0.9 New Taiwan dollar or US$0.03, with floating value per point under the global budget scheme) as a case management fee per participant per year, along with a 550 points bonus if the performance of their CHCG reaches a specified quality indicator goal (online supplemental appendix 1). The cost of FPICP is relatively low compared with its coverage rate. It requires a share of just 0.2% (US$40 million) of Taiwan’s US$20 billion NHI annual budget.

The impact of the FPICP since its reformation in 2010 has yet to be fully ascertained. Therefore, we aimed to quantify the progress of the FPICP towards the NHI administration’s goal of fortifying primary healthcare and reducing wastage of medical resources. One indicator of choice was the hospitalisation rate for ambulatory care sensitive conditions (ACSCs), which are considered manageable by primary care physicians and as such hospital stays from ACSC can be considered preventable. We hypothesised that patients participating in the FPICP would have a lower rate of hospitalisation for ACSC compared with non-participating patients.

**Methods**

We conducted a population-based cohort study comparing hospitalisation rates for ACSC among FPICP participating and non-participating patients in Taiwan. The study was in compliance with the Strengthening the Reporting of Observational Studies in Epidemiology checklist of essential items (version 4) for cohort studies.

**Setting**

The study used data from the FPICP database, the reimbursement database of Taiwan’s NHI administration containing all NHI administration-selected patients for FPICP enrolment. The data of patients registered in FPICP in fiscal years 2011–2015 were extracted, and the follow-up time interval to calculate the rate of hospitalisation was the 1 year in which the patient was selected for FPICP enrolment.

The consulted database has a data structure and format similar to the main NHI administration reimbursement database and the National Health Insurance Research Database, on which 99.6% of Taiwan’s population are enrolled. The database also contained comprehensive drug prescription files and original claim data. In addition to the main NHI administration reimbursement database, the database of the FPICP included four other components: a dataset on the original FPICP candidates, a dataset on the final FPICP participants, a dataset on CHCG profiles and a dataset on quality assessments of the CHCG. The database of the FPICP, with the help of these extra datasets, enables research specific to the family physician system in Taiwan and was first used in a recent publication by the authors.

**Target population**

The study’s target population was patients in primary care clinics eligible for FPICP inclusion in the fiscal year 2011–2015 and aged above 5 years. The original version of the ACSC in the pan-Canadian primary healthcare indicators targeted patients aged 5–75 years for the calculation of hospitalisation rates. However, because the FPICP focused on the elderly and patients aged over 80 accounted for 6%–9% of all FPICP participants, we did not apply an upper age limit for our target population.

**Variables**

When defining ACSC in this study, we referred to the standards set by the Canadian Institute for Health Information and modified them according to primary healthcare practice routines in Taiwan. The outcome measures were the rates of hospitalisation for ACSC including asthma/chronic obstructive pulmonary disease (COPD), diabetes or its complications and heart failure. Rate of hospitalisation due to ACSC was calculated as the number of patients hospitalised with a main discharge diagnosis of ACSC per 1000 of the outpatient population with ACSC. Specifically, the numerator is hospitalisation with a main diagnosis of one of the conditions below:
1. COPD/asthma: ICD-9-CM codes that begin with 490–496; 480–488, and with a secondary diagnosis 490–496; or ICD-10-CM codes that begin with J10.0–J18, or J20–J22, and with a secondary diagnosis J40–J47.

2. Diabetes and its complications: ICD-9-CM codes that begin with 250; or ICD-10-CM codes that begin with E10, E11, E13.

3. Heart failure: ICD-9-CM codes that begin with 428 or 518.4; or ICD-10-CM codes that begin with I50 or J81.

The denominator is the population with an outpatient diagnosis of ACSC in the previous year.

The intervention of interest was enrolment in the FPICP or not (figure 1). These NHI Administration-selected candidates became FPICP participants if their primary care physicians joined the project and enrolled all assigned patients as participants. Factors regarded as potential confounders included age, sex, monthly income, region of residence and comorbidities. Monthly income and region of residence were based on the Registry for Beneficiaries dataset obtained on a study subject’s enrolment in the NHI programme. Comorbidities were assessed using the Charlson Comorbidity Index (CCI). We defined the diagnosis of a comorbidity as receiving the same diagnosis no less than twice in the previous year based on the ICD-9-CM and ICD-10-CM codes as indicated by physicians’ claims data. The ICD codes used in this study are described in online supplemental appendix 2. The technical part of CCI calculation was based on the open-sourced SAS scripts published by Healthcare Delivery Research at the National Cancer Institute of the USA.

We converted the quantitative variables into categorical ones as follows: older adults were defined as participants aged 65 years or older, which is consistent with the WHO’s definition; monthly income was categorised by tertiles; codes of residential region were transformed into three levels of urbanisation according to Taiwan National Health Research Institute (NHRI) publications, with level 1 referring to ‘most urbanised’ and level 3 ‘least urbanised’ communities; increased comorbidity score was defined as a CCI of 3 or greater, which was adopted or suggested by previous studies.

### Statistical analysis

Values were presented either as percentages or as arithmetic means with SD in descriptive analyses. Logistic regression was used to calculate the ORs for the influence of FPICP participation on the rate of hospitalisation for ACSC. Age (in dichotomised categories by 65 years old), gender and level of comorbidities were included as independent variables in the model. A two-tailed p value of 0.05 was considered statistically significant, and 95% CIs were also calculated. Propensity score matching for FPICP participating and non-participating patients was not applied due to the large number of observations (1–2 million participants per year) and limited computing power. All statistical analyses were conducted using SAS software (V.9.4SAS Institute).

### Patient and public involvement

We did not directly include patient and public involvement in this study, but the database used in the study was developed with patient and public involvement and is updated by a committee that includes patient representatives from the NHI Administration, Ministry of Health and Welfare, Taiwan.

### RESULTS

After excluding children under the age of five and patients who had dropped out from the NHI programme before recruitment (deceased or moved away), the study population, including participants and non-participants, was 3.94 million in 2011 and 5.34 million in 2015 (online supplemental appendix 3). Among them, the population of FPICP participants was 2,316,114 (43.4%) for 2015.

Table 1 summarises the demographic characteristics of the study participants in 2015. For FPICP participants,
was 366 with ACSC (COPD/asthma, diabetes or heart failure)

and for diabetes or its complications (12.7‰–18.1‰ vs

for COPD/asthma (37.9‰–42.3‰ vs 28.6‰–35.9‰)

participants, FPICP participants had lower hospitalisation

supplemental appendix 4).

19.3%). (full comparison from 2011 to 2015 in online

ment (OPD)/clinic visits (by 9.0%–15.2%), for ER visits

by 11.4%–14.5%) and for hospitalisation (by 17.5%–

a smaller proportion of patients with CCI >2 (by 1.4%)

and 20.5% were aged under 20. As differences were small

53.6% of them were female, 17.3% were aged over 70

and our knowledge, this study is the first to directly use

real-world data from the FPICP to verify the effectiveness

of the programme. Moreover the reason for reporting

the outcome of ACSC is that the quality of care for these

diseases can be reflected in a reduction in the use of

hospital resources if well controlled at primary healthcare

316 114 3 021 263

Table 1 Characteristics of study base at enrolment in 2015

FPICP Non-FPICP

| Number of observation | 2 316 114 | 3 021 263 |
|-----------------------|-----------|-----------|
| Sex                   |           |           |
| Female                | 1 241 437 (53.6%) | 1 613 354 (53.4%) |
| Male                  | 1 074 677 (46.4%) | 1 407 908 (46.6%) |
| Age (years)*          |           |           |
| 5–10                  | 222 703 (9.6%) | 239 933 (7.9%) |
| 10–20                 | 252 397 (10.9%) | 303 073 (10.0%) |
| 20–30                 | 155 892 (6.7%) | 161 008 (5.3%) |
| 30–40                 | 259 820 (11.2%) | 287 288 (9.5%) |
| 40–50                 | 291 989 (12.6%) | 344 114 (11.4%) |
| 50–60                 | 381 070 (16.5%) | 486 180 (16.1%) |
| 60–70                 | 351 176 (15.2%) | 524 064 (17.3%) |
| 70–80                 | 249 923 (10.8%) | 423 040 (14.0%) |
| 80–90                 | 131 147 (5.7%) | 217 834 (7.2%) |
| over 90               | 19 797 (0.8%) | 34 728 (1.3%) |
| Monthly income†‡      |           |           |
| Level 1 (high)        | 826 853 (35.7%) | 1 078 591 (35.7%) |
| Level 2 (medium)      | 817 588 (35.3%) | 1 102 761 (36.5%) |
| Level 3 (low)         | 671 673 (29.0%) | 839 910 (27.8%) |
| Urbanisation‡         |           |           |
| Level 1 (high)        | 528 074 (22.8%) | 797 613 (26.4%) |
| Level 2 (medium)      | 1 100 154 (47.5%) | 1 326 334 (43.9%) |
| Level 3 (low)         | 687 886 (29.7%) | 897 315 (29.7%) |

*Age at enrolment.
†Counted in New Taiwan dollar (NTD).
‡Categorised by tertiles.

FPICP, Taiwan’s Family Practice Integrated Care Project; .

53.6% of them were female, 17.3% were aged over 70
and 20.5% were aged under 20. As differences were small
between FPICP participants and non-participants in terms
of monthly income (29.0% vs 27.8% in the low-income
category) and urbanisation level of their residential area
(29.7% vs 29.7% in the low-urbanisation category), we
did not include monthly income and the urbanisation as
independent variables in the logistic regression analysis.

Table 2 shows that the outpatient population of patients
with ACSC (COPD/asthma, diabetes or heart failure) was
366 047 among FPICP participants and 481 600 among
non-participants in 2015. FPICP participants had a smaller
proportion of patients with CCI >2 (by 1.4%), reduced medical
costs per year for outpatient department (OPD)/clinic visits
(by 9.0%–15.2%), for ER visits (by 11.4%–14.5%) and for hospitalisation (by 17.5%–
19.3%). (full comparison from 2011 to 2015 in online supplemental appendix 4).

Figure 2 shows the rate of hospitalisation for selected
ACSCs from 2011 to 2015. Compared with non-
participants, FPICP participants had lower hospitalisation
for COPD/asthma (37.9‰–42.3‰ vs 28.6‰–35.9‰)
and for diabetes or its complications (12.7‰–18.1‰ vs
10.8‰–14.9‰) (p<0.05). The reduced hospitalisation
rate for heart failure was also noted, but there was no sta-

tistical significance (49.6‰–54.1‰ vs 43.9‰–50.6‰).

After adjusting for age, sex and level of comorbidities by
conditional logistic regression, participation in the FPICP
was associated with lower hospitalisation for COPD/
asthma (OR 0.91, 95% CI 0.87 to 0.94) and for diabetes or
its complications (OR 0.87, 95% CI 0.83 to 0.92) but not
for heart failure (OR 0.97, 95% CI 0.88 to 1.07) (table 3).

DISCUSSIONS
Main findings
The FPICP has been the most important reform
programme for primary healthcare in Taiwan since 2010,
and to our knowledge, this study is the first to directly use
real-world data from the FPICP to verify the effectiveness
of the programme. Moreover, the reason for reporting
the outcome of ACSC is that the quality of care for these
diseases can be reflected in a reduction in the use of
hospital resources if well controlled at primary healthcare
clinics. We found that participants in the FPICP presented
a lower hospitalisation rate regarding ACSC, including
asthma/COPD and diabetes or its complications. After
adjusting for variables such as age, sex and comorbid-
ities, participation in the FPICP remains an independent
protective factor for preventable hospitalisation. This
major finding sheds light on the team-based primary
healthcare model such as FPICP strengthened primary

Table 2 Comorbidities and utilisation of medical resource among patients with ACSC, by FPICP participation (2015)

| FPICP            | Non-FPICP       |
|------------------|-----------------|
| Number of observation | 366 047          | 481 600          |
| CCI               |                 |
| High (>2)         | 52 656 (14.4%)  | 76 019 (15.8%)  |
| Low (0–2)         | 313 391 (85.6%) | 405 581 (84.2%) |
| Clinic/outpatient care |             |
| Number of visits/year | 12.6 (12.1) | 14.0 (13.3) |
| Medical cost/year (point)* | 9458 (67 589) | 10 655 (34 250) |
| Emergency care    |                 |
| Number of visits/year | 0.43 (1.49) | 0.45 (1.44) |
| Medical cost/year (point) | 1365 (5670) | 1541 (7400) |
| Inpatient care    |                 |
| Number of visits/year | 0.30 (0.77) | 0.35 (0.85) |
| Medical cost/year (point) | 18 341 (91 482) | 22 733 (96 611) |
| Hospitalisation rate | 16.3%          | 18.6%           |
| Length of stay (day) | 16.7 (39.4) | 17.7 (35.5) |

SD or percentage is shown in parentheses.
*Floating point value (1 point ~NT$0.9) under global budget scheme since 2001.
ACSC, ambulatory care sensitive conditions; CCI, Charlson Comorbidity Index; FPICP, Taiwan’s Family Practice Integrated Care Project.
healthcare capacity and improved quality of community healthcare.  

Other than the above findings, FPICP participants were also found to have lower medical costs as outpatients, through emergencies and through hospitalisation, compared with the non-participants. The difference in hospitalisation costs is particularly significant (17.5%–19.3%), followed by emergency costs (11.4%–14.5%) and outpatient expenses (by 17.5%–19.3%).

Our study supported the evidence that by engaging in data-driven, continuous quality improvement, team-based care can offer higher accessibility to care, as well as more effective and efficient delivery by providing care coordination. 17–20 In Canada, Carter et al found moderate quality evidence that team-based models of care led to reductions in emergency department use, but the evidence was mixed for hospital admissions. 21 McAlister et al also demonstrated that care within a primary care network was associated with fewer emergency department visits and fewer hospital days. 22

Regarding disease-specific team-based care examples, a meta-analysis done by Carter et al demonstrated that team-based care was associated with improved blood pressure control. 23 Proper training, the use of an electronic clinical reminder system and the enhanced engagement of registered nurses can help to improve completion rates of asthma action plans in a team-based primary care setting. 24 As to diabetes care, team-based care management interventions that use nurses, medical assistant health coaches, and behavioural specialists to support diabetes patients can help primary care practices achieve value-based targets of improved health, cost and patient experience. 25,26 Furthermore, in patients with COPD, a team-based approach following the treatment guidelines of Global Initiative for Chronic Obstructive Lung Disease is critical to successfully implement comprehensive care. 26

As to hospitalisations for heart failure, the lack of a significant difference between enrolled and unenrolled participants was observed from 2011 to 2015, except in 2012. There are several possible explanations. First, a lack of close cooperation with heart failure care teams and low-intensity transitional care may have contributed to the limited efficacy. Treatment for heart failure patients in Taiwan is usually referred to cardiologists rather than follow-up at community-based clinics because diagnostic procedures need to be done such as echocardiograms or interventional studies in Taiwan. Second, the hospitalisation rate for enrolled persons with heart failure was indeed lower than non-enrolled across 2011–2015 although there was no statistical difference. To be noted, the cases numbers for heart failure hospitalisation were smaller compared with diabetes or COPD cases (figure 2, online supplemental appendix 5–7). As larger the number of cases, the higher the statistical power, and more likely the difference to be significant. Moreover, referring patients to specialised outpatient heart failure clinics, staffed with trained healthcare providers who are familiar with current guidelines and available resources, has been shown to reduce hospital admissions. 27–29 High- or moderate-intensity transitional care
interventions combining home visits with follow-up telephone calls, clinic visits or both reduced readmission risk if implemented for a longer period, for example, at least 6 months. One Cochrane review from Takeda et al in 2019 found low quality of evidence that multidisciplinary interventions instead of clinic-based interventions may reduce the risk of readmission for heart failure. Variations in study location and time of occurrence hamper attempts to review costs and cost-effectiveness.

What Taiwan FPICP highlights is on integrated care. Primary care physicians of FPICP regularly share case discussions with medical specialists in the hospitals, so the care for chronic diseases might be more in line with current clinical guidelines. These better medical cares are reflected in lower hospitalisation rates among patients with diabetes and COPD; however, for patients with heart failure, shared care by cardiologists in the hospitals is often needed and sometimes the patients may require planned hospitalisation to do interventions; therefore, the advantages of FPICP in primary care system are more difficult to be reflected. This may help explain why there might not have been a significant difference in hospitalisations between enrolees and non-enrollees with heart failure.

Above all, FPICP as a team-based care model encourages community clinics and hospitals to form cooperative networks to facilitate the improvement of quality care, through data-driven, continuous care and coordination.

Policy implications

The quality assessment of the FPICP involves processes of care which are primarily preventive healthcare services, such as influenza vaccinations, adult regular health examinations and nationwide cancer screenings including Pap smear, fecal immunochemical tests and oral cancer screening by inspection. The ACSC hospitalisation rate highlighted in this study is only approximately one-tenth of all quality assessment items for the FPICP (online supplemental appendix 1). It is worth noting that the FPICP assesses quality indicators in a group-wise manner, scoring each CHCG instead of individual physicians. The rewards such as bonus payments and options to include more patients were also granted based on the performance of the CHCG. Physicians within a CHCG work as a group to review their performance and facilitate two-way coordination with their backup hospitals through regular monthly meetings. This healthcare model may be one of the keys to improving the overall quality of care in the community.

Except for copayments or out-of-pocket expense differences, whether patients are confident of their diseases being handled in community clinics is a crucial factor in forming effective CHCGs to achieve universal health coverage. If a community clinic is not capable of providing care for nearby patients who fail to seek medical help in outpatient departments of hospitals due to expense issues, the long-term consequence is likely to increase medical costs in terms of emergencies and hospitalisation. While reducing medical expenses in various ways is imperative for policymakers, patients care about the quality and accessibility of medical care. Minimising preventable hospitalisations by strengthening the ambulatory care capability of primary clinics is one of a few approaches that improves medical quality while maintaining or even reducing overall medical expenses, satisfying both patients and payers of the healthcare system. Potentially, an effective programme such as the FPICP might help enhance patients’ trust in their family doctors and decrease unnecessary emergency visits or hospitalisations.

Limitations

Similar to previous database studies using physician claim data, our research, which was based on reimbursement databases, also has some limitations. First, we did not acquire data regarding patients’ diets, physical activities, alcohol/cigarette consumption, etc., and these potential confounding factors may affect clinical outcomes. Second, we were not able to apply propensity score matching techniques due to limitations in computing power for the huge amount of data acquired, although the FPICP participating and non-participating patients were mostly assigned by NHI administration based on the same criteria. Nonetheless, we applied multiple logistic regression to adjust for potential differences in demographics and comorbidities. Third, our research only determined whether an association existed between the FPICP and the outcomes of interest, rather than causality. If an association was significant, it could be that the FPICP led to better outcomes or that physicians with better clinical ability were more likely to join the FPICP and be rewarded.

In conclusion, the FPICP is a team-based care model and a modified P4P programme. It features mandatory inclusion of NHI administration-assigned patients with high medical needs in ambulatory care, and operates through a CHCG formed by local clinics, is vertically coordinated with regional hospitals. Our study adopted a population-based cohort design to validate the effectiveness of this model and found that participation in the FPICP is an independent protective factor for preventable hospitalisation. The observed trend also showed lower overall medical costs in FPICP participating patients. The experience of the FPICP may serve as a reference for policymakers in developing primary care reform programmes in order to achieve universal health coverage and improve the quality of community healthcare.

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Patient consent for publication Not required.

Ethics approval This article does not refer to any studies conducted by the authors on human or animal subjects. This study was approved by the ethical review board of the National Taiwan University Hospital. (NTUH2017I1086RIND).

Data availability statement Data may be obtained from a third party and are not publicly available. The data support the findings of this study are available from National Health Insurance Administration (Taiwan), but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of National Health Insurance Administration (Taiwan).

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