Evaluation of pharmacoinvasive strategy versus percutaneous coronary intervention in patients with acute myocardial infarction with ST-segment elevation at the National Institute of Cardiology (PHASE-MX)

Evaluación de la estrategia farmacoinvasiva versus angioplastia coronaria transluminal percutánea primaria en pacientes con infarto agudo de miocardio con elevación del segmento ST en el Instituto Nacional de Cardiología (PHASE-MX)


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Abstract

Objective: The objective of PHASE-MX registry is to validate the efficacy and safety of the pharmacoinvasive strategy in comparison with percutaneous coronary intervention (PCI) in patients with acute myocardial infarction with ST segment elevation (STEMI) in a metropolitan region of Mexico. The primary outcome will consist of the composite of cardiovascular death, re-infarction, stroke and cardiogenic shock. Methods: The PHASE-MX registry will include a prospective cohort of patients with STEMI who received reperfusion treatment (mechanical or pharmacological) in the first 12 h after the onset of symptoms. The registry is designed to compare the efficacy and safety of primary PCI and pharmacoinvasive strategy. The simple size was calculated in 344 patients divided into two groups, with an estimated loss rate of 10%. Patients included in the PHASE-MX cohort will be followed for up to one year. Conclusion: In Mexico, only 5 out of 10 patients with STEMI have access to reperfusion therapy. Pharmacoinvasive strategy is takes advantage of the accessibility of fibrinolysis and the effectiveness of PCI. The present research protocol aims to provide information that serves as a link between information derived from controlled clinical trials and records derived from real world experience.

Key words: Acute myocardial infarction. Pharmacoinvasive. Reperfusion. Mexico.

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Resumen

Objetivo: El objetivo del registro PHASE-MX es validar la eficacia y seguridad de la estrategia farmacoinvasiva en comparación con la angioplastia coronaria transluminal percutánea primaria (ACTPp) en pacientes con infarto agudo de miocardio con elevación del segmento ST (IAMCEST) en una región metropolitana de México. El desenlace primario es el compuesto de muerte cardiovascular, reinfarto, accidente vascular cerebral y choque cardiogénico. Métodos: El registro PHASE-MX es una cohorte prospectiva de pacientes con IAMCEST que recibieron tratamiento de reperfusión (mecánico o farmacológico) en las primeras 12 horas desde el inicio de los síntomas, atendidos en el Instituto Nacional de Cardiología Ignacio Chávez. El análisis estadístico se basa en la no inferioridad de la estrategia farmacoinvasiva en comparación con la ACTPp. Se calcula un tamaño de muestra de 344 pacientes divididos en dos grupos (angioplastia primaria y estrategia farmacoinvasiva), considerada una tasa de pérdidas de 10%. Los pacientes incluidos en la cohorte PHASE-MX se seguirán durante un año. Discusión: En México, sólo 5 de cada 10 pacientes con IAMCEST tienen acceso al tratamiento de reperfusión. La estrategia farmacoinvasiva aprovecha la accesibilidad de la fibrinólisis y la efectividad de la ACTPp, por lo que podría resultar el método de elección en el tratamiento del IAMCEST en la mayoría de los casos. El presente protocolo de investigación pretende aportar información que sirva como enlace entre la información derivada de los estudios clínicos controlados y los registros derivados de la experiencia del mundo real.


Introduction

Mexico is the country with the highest mortality rate of ST-segment elevation myocardial infarction (STEMI) among the Organization for Economic Cooperation and Development (OECD) member countries, with an estimate of 26.6/100 discharges, in comparison with OECD average of 8 deaths/100 discharges.

Reperfusion treatment in STEMI has been shown to decrease long-term morbidity and mortality. The European Society of Cardiology clinical practice guidelines recommend the use of fibrinolysis or primary percutaneous coronary intervention (PCI) with a Class I recommendation and Grade A evidence, within the first 12 h of symptom onset. Although pPTCA is superior to fibrinolysis when prospectively compared in randomized controlled trials, real-life registries suggest that fibrinolysis is an efficacious treatment when timely administered and that it prevents delays related to PCI.

The pharmacoinvasive strategy consists of immediate fibrinolysis and subsequent systematic coronary angiography, preferably within the first 3-24 h after fibrinolysis. The advantage of this strategy allows easy access to fibrinolytic combined with percutaneous transluminal coronary angioplasty efficacy to prevent reinfarction and recurrent ischemia. The pharmacoinvasive strategy has been shown not to be inferior in prospective clinical studies and real-life registries, and multiple countries have published their experience with regard to pharmacoinvasive systems.

The pharmacoinvasive strategy could be a feasible approach in countries and regions where access to PCI is limited due to geographic, cultural, or social reasons, such as Mexico. However, the pharmacoinvasive strategy has not been prospectively validated in Mexican patients with acute STEMI.

The purpose of the PHASE-MX study is to validate the efficacy and safety of the pharmacoinvasive strategy in comparison with primary angioplasty in patients with acute STEMI in a metropolitan region of Mexico.

Methodology

PHASE-MX is a prospective cohort study of patients with ST-segment elevation acute myocardial infarction treated with fibrinolysis or primary angioplasty and finally attended in a tertiary care center in Mexico City. The project is an initiative of the Emergency and Coronary Care Unit of the Ignacio Chávez National Institute of Cardiology and consists of a prospective record of all STEMI cases that received reperfusion treatment within the first 12 h. Follow-up will be extended for up to 12 months after hospital discharge.

Study population

Patients older than 18 years with a STEMI diagnosis, attended in any of the hospitals belonging to the metropolitan area of Mexico City, in the process of reperfusion treatment (pharmacological or mechanical) within the first 12 hours of symptom onset and referred to the Ignacio Chávez National Institute of Cardiology. The area of care and reference of the Mexico City metropolitan area comprise a calculated area of 7954 km², with a population of 20.4 million inhabitants (Fig. 1 and Table 1). It will not include patients whose diagnosis at hospital discharge is other than acute STEMI.
Objectives

The general purpose of the PHASE-MX registry will be to assess the efficacy of the pharmacoinvasive strategy in comparison with primary percutaneous transluminal coronary angioplasty in patients with ST-segment elevation myocardial infarction in the primary combined outcome of cardiovascular death, reinfarction, stroke, and cardiogenic shock at in-hospital follow-up. Specific objectives will include knowing the frequency of major bleeding, stroke, and all-cause mortality at 12 months of follow-up.

Sample size calculation

The primary objective will be to compare the composite outcome of cerebrovascular disease (CVD), reinfarction, cardiogenic shock, and cardiovascular death, after predicting a statistical power (1-β) of 80%, an α level of 0.05, and taking into account, a predicted rate of the primary outcome of 12.4% at follow-up day 30 in patients treated with reperfusion and a maximum tolerated degree of difference (non-inferiority – d) of 5%; a total sample of 157 patients per arm are calculated for demonstrating non-inferiority of the pharmacoinvasive strategy versus primary angioplasty. If an attrition rate of 10% is assumed, a final total sample of 344 patients is calculated. Enrollment started on April 1, 2018, and its completion is expected by March 31, 2020. The review and analysis of the information are expected to be available to submit for publication to a journal of large circulation by the second semester of 2020.

Follow-up

From the moment of inclusion, clinical follow-up will be carried out until hospital discharge. Then, upon completing 12 months of follow-up (accounted for since hospital discharge), vital state will be analyzed by means of telephone contact.

Table 1. Hospitals by state of the metropolitan area of Mexico city that referred patients with STEMI to INCICH

<table>
<thead>
<tr>
<th>Localization of hospitals that refer to INCICH</th>
<th>Average distance to INCICH (km) (shortest-longest)</th>
<th>Average transportation time (minutes)** (shortest-longest)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mexico city (15)</td>
<td>21.6 (12.5-31.8)</td>
<td>46.9 (18-100)</td>
</tr>
<tr>
<td>National Health Institutes** (5)</td>
<td>1.9 (0.5-6.6)</td>
<td>6 (3-11)</td>
</tr>
<tr>
<td>State of Mexico (14)</td>
<td>88.8 (29.6-190)</td>
<td>91.8 (30-150)</td>
</tr>
<tr>
<td>Total (34)</td>
<td>37.4</td>
<td>48.2</td>
</tr>
</tbody>
</table>

INCICh (Instituto Nacional de Cardiología Ignacio Chávez): Ignacio Chávez National Institute of Cardiology.

*Time calculated using the Google Maps app at 12:00-14:00 h, Mexico City time zone.

**Includes the National Institute of Medical Sciences and Nutrition Salvador Zubirán, General Hospital Dr. Manuel Gea González, National Institute of Respiratory Diseases Ismael Cossio Villegas, National Institute of Cancer, National Institute of Neurology and Neurosurgery Dr. Manuel Velasco.

Figure 1. Satellite map showing the area served by the Ignacio Chávez National Institute of Cardiology (INCICh – Instituto Nacional de Cardiología Ignacio Chávez). Blue indicators point at the localization of hospitals where patients with STEMI received reperfusion treatment with pharmacological fibrinolysis and then were referred to the INCICH (marked on the map with a white cross on red background), for a period of approximately 1 year. The Valley of Mexico metropolitan area and the area served by INCICH have an extension of approximately 8000 and 36,000 km², respectively.
Statistical Analysis

Study variables

In addition to the main variables, the following will be collected:
- **ST-segment elevation myocardial infarction**: patients with chest pain of angina pectoris with ST-segment elevation on 12-lead electrocardiogram. ST-segment elevation should appear on two contiguous leads, larger than 2.5 mm in males younger than 40 years, larger than 2 mm in males older than 40 years, or larger than 1.5 mm in females, on V2-V3 electrocardiographic leads or larger than 1 mm on any other lead.
- **Time to first medical contact**: period of time, measured in minutes, from the onset of symptoms indicative of ischemia until receiving medical care for the first time. This can be provided by a health professional or medical emergency technician.
- **Site of first medical contact**: place where initial medical care was provided. It may correspond to any primary, secondary, or tertiary medical center.
- **Door-to-needle time**: period of time, measured in minutes, from the moment of entering the site of first medical contact to the moment of receiving the fibrinolytic drug.
- **Door-to-balloon time**: period of time, measured in minutes, from the moment of entering the health-care site with PCI capacity to the advancement of the metal guidewire beyond the lesion in the infarct-related artery (IRA).
- **Total ischemia time**: time elapsed, measured in minutes, from the onset of symptoms indicative of ischemia to the moment of advancement of the metal guidewire beyond the lesion in the IRA. In case of receiving fibrinolysis, it will be measured from the onset of symptoms until evidence of reperfusion on ECG, after 90 min of fibrinolytic treatment administration.

Effectiveness criteria

Angiographic: TIMI Grade 3 anterograde flow in IRA and TMP 3 myocardial perfusion.

Electrocardiographic: ST-segment decrease of at least 50% versus initial electrocardiogram, 90 min after fibrinolytic agent administration.

Primary outcomes


Secondary outcomes

- **Major hemorrhage**: hemorrhage leading to death, retroperitoneal bleeding, central nervous system bleeding (including the eye), bleeding with hemodynamic repercussion, bleeding that requires surgical intervention, decompression of a closed compartment, transfusion of one or more units of packed red blood cells, hemoglobin decrease greater than 3 g/dL, or hematocrit decrease greater than 10% with regard to a previous determination.
- **Minor hemorrhage**: microscopic hematuria unrelated to urethral trauma, epistaxis requiring tamponade or surgical intervention, gastrointestinal tract bleeding, conjunctival bleeding, hematoma larger than 5 cm, bleeding reducing hemoglobin less than 3 g/dL.
- **TIMI flows**, documented by invasive coronary angiography, and categorized as follows: TIMI 0: complete flow absence; TIMI 1: contrast medium partial penetration, without crossing the causative lesion or filling the distal coronary artery; TIMI 2: contrast medium penetration beyond the causative lesion, without filling the coronary artery or, otherwise, slower than normal coronary arteries; and TIMI 3: normal coronary arterial flow.
- Mortality at 12 months of follow-up.

Discussion

The PHASE-MX study aims to demonstrate, through a real-life observational analysis (i.e., without randomization or control of variables), that fibrinolysis and subsequent PCI in patients with STEMI is not inferior in efficacy versus PCI. The strengths of this study lie on three points; the first one refers to the absence of randomization and control groups; the second, to fibrinolysis administration in a wide variety of clinical contexts, and the third, to the performance of PCI in a national reference center, regarded as high volume in coronary care.
angioplasty. The main weaknesses of this study include the probability of patients not being transferred to the inclusion center due to the severity of their clinical condition and the wide variety of coadjuvant treatment.

One of the main implications derived from the findings of this study is to support a wider use of fibrinolysis in Mexico, where availability of hemodynamic laboratories is limited. In addition, it is likely to be the scientific support of promotional campaigns for the care of myocardial infarction in Mexico.

Funding

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Conflicts of interest

The authors declare that they have no conflicts of interest.

Ethical disclosures

Protection of people and animals. The authors declare that no experiments were performed on humans or animals for this investigation.

Confidentiality of data. The authors declare that they have followed the protocols of their work center on the publication of patient data.

Right to privacy and informed consent. The authors declare that no patient data appear in this article.

This research protocol was submitted to the Committee of Ethics and Research of Instituto Nacional de Cardiología Ignacio Chávez, with registration number PT-19-109. In addition, it was registered on the ClinicalTrials.gov platform with identifier NCT03974581.

References