A proposed functional clinical classification predicts in-hospital and long-term survival in the setting of acute right ventricular infarction

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Summary

Background: The objectives of the present investigation were to validate the prognostic role of a proposed Clinical Classification [CC], to evaluate the TIMI risk score [RS] and to establish whether the TIMI-RS should incorporate points for patients with acute right ventricular infarction [TIMI-RS-RVI]. Methods and results: A total of 523 RVI patients were classified on clinical and functional basis as: A, without right ventricular failure [RVF], B with RVF and C with cardiogenic shock. The CC was evaluated prospectively among 98 patients with RVI and retrospectively in 425 RVI patients. The TIMI-RS was evaluated prospectively among 622 patients with STEMI [anterior:277, inferior:247, RVI:98], and retrospectively in 425 RVI patients. The CC established differences among the 3-RVI Classes for in-hospital mortality [prospectively and retrospectively; p < 0.01, p < 0.001, respectively] that were maintained at 8 years [p < 0.001]. Patients with anterior and inferior STEMI, but not those with RVI revealed an association between outcome and TIMI-RS [p < 0.001]. Testing for TIMI-RS-RVI did not result a good prognostic tool [ROC = 0.9; excellent discrimination, but with a very poor “clinical calibration”]. Conclusions: The proposed CC allowed prediction of mortality at short- and long-term in the setting of acute RVI. The role of the TIMI-RS should be reevaluated prospectively.

Resumen

PROPUESTA DE UNA CLASIFICACIÓN FUNCIONAL Y CLÍNICA PARA PREDETERMINAR LA SOBREVIDA INTRAHOSPITALARIA Y A LARGO PLAZO EN ENFERMOS CON INFARTO AGUDO DEL VENTRÍCULO DERECHO

Antecedentes: Esta investigación tuvo como metas validar el significado pronóstico de la clasificación clínica [CC] previamente propuesta, valorar el puntaje TIMI y tratar de establecer si es posible incorporar puntos clínicos/ electrocardiográficos al puntaje TIMI en enfermos con infarto agudo del ventrículo derecho [TIMI-IAVD]. Métodos y resultados: Un total de 523 enfermos con IAVD se clasificaron con fundamento funcional y clínico en: Clase A, sin disfunción ventricular derecha [DVD], Clase B con DVD y en Clase C en estado de choque. La CC se sustentó con bases ecocardiográficas y hemodinámicas. Fue evaluada de manera prospectiva en 98 enfermos y de forma retrospectiva en 425 enfermos con IAVD. El puntaje TIMI fue valorado de manera prospectiva en 622 enfermos con SICA con elevación del segmento ST [anteriores: 277, inferiores: 247 y con IAVD: 98] y de forma retrospectiva en 425 con IAVD. La CC estableció diferencias en relación a la mortalidad intrahospitalaria en los tres grupos de enfermos [de manera prospectiva y retrospectiva; p < 0.01, p < 0.001, respectivamente], las que se mantuvieron durante 8 años de seguimiento [p < 0.001].

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as a prognostic tool in the scenario of RVI patients.

**Conclusiones:** La CC propuesta permite predecir la mortalidad a corto y a largo plazo en el escenario del IAVD. Debe revalorarse la utilidad del puntaje TIMI para el IAVD de forma prospectiva y en una cohorte más amplia. (Arch Cardiol Mex 2008; 78: 369-378)

**Key words:** Right ventricular infarction. TIMI risk score. Clinical classification.

**Introduction**

The development of the TIMI [Thrombolysis in Myocardial Infarction]-risk score [RS] has proved to be a useful tool to easily stratify patients with ST-elevation myocardial infarction [STEMI] and its usefulness has also been validated in a large STEMI registry. While several investigators have established the prognostic role of right ventricular infarction [RVI] in patients, only few have previously attempted to stratify risk among patients with RVI for in-hospital and long-term survival. Therefore, further studies are required to better define the prognostic importance of right-sided ST-elevation and to establish whether the TIMI-RS should incorporate points for right-sided electrocardiographic changes. Furthermore, the pathophysiologic mechanism of low output in RVI is quite different from that of left ventricular infarction. Thus the criteria for inclusion in clinical classes of left ventricular STEMI can not be used for RVI. In a previous study we demonstrate that there is a wide “clinical spectrum of acute RVI”, that three classes could be identified prospectively and that there are significant differences in early mortality rates between RVI classes. Accordingly, if RVI is documented, it must then be stratified using the clinical classification as has been proposed, because Class A RVI patients, those without right ventricular failure [RVF] was not found to be a predictor of mortality. Class B [with RVF] and class C [with cardiogenic shock] are strongest independent predictors of in-hospital mortality. Therefore, we recommend that the “spectrum of RVI” should always be considered in discussion clinical scenario of RVI.

**Methods**

On the previous mentioned background, the aims of this study were: First, to ascertain the role of a proposed clinical classification [CC] in the prediction of short-term mortality and long-term survival for RVI patients. Second, to evaluate the TIMI-RS and if incorporating points for clinical RVI findings and right-sided electrocardiographic changes [TIMI-RS-RVI] could identify better the prognosis in patients with acute RVI.

**Patient population**

The study protocol was approved by the ethics review commission. We prospectively screened 1,402 consecutive patients admitted with a first inferior STEMI from 1990 to 2002, and identified 425 with extension to the right ventricle [Right ventricular infarction: RVI]. This RVI population underwent follow-up through July 2007. Patients were included in the study if they all had a complete clinical examination, an echocardiogram, or a coronary angiogram or a right heart catheterization. Isolated acute RVI, history of valve heart disease, previous heart failure, pulmonary arterial hypertension or renal failure were exclusion criteria for the enrollment to the study. We did not include in the shock category patients with hypotension related to hypovolemia, transient hypotension related to vasodilatation and bradycardia associated with spontaneous reperfusion [Bezold-Jarisch reflex], or hypotension related to atrioventricular heart block or cardiac arrhythmias or associated with mechanical complications [ventricular septal or myocardial wall rupture or cardiac tamponade] at entry.
Diagnosis for RVI, RVF and cardiogenic shock

The diagnostic criteria for inferior myocardial infarction with extension to the walls of the right ventricle, RVF and cardiogenic shock have been published previously. Briefly, in addition to standard ECG leads, right-sided thoracic ECG [leads V3R-V7R] was recorded in all patients immediately after admission. ST-segment elevation in lead V4R of greater than 0.1mV was used to diagnose RVI. The diagnosis of RVI was also based on clinical features that have been described previously as being associated with this variety of infarction, and also on echocardiographic findings. Echocardiograms were analyzed separately by two skilled cardiologists who were blinded to the clinical characteristics, ECG findings, angiographic and hemodynamic information of RVI patients. RVF diagnosis, was based on clinical features [persistently systemic hypotension [systemic systolic pressure [SSP] ≤ 100 mm Hg, right-sided S3 and S4]] but without features of shock; echocardiographic evidence of ischemic RVF [for example RV wall motion abnormalities [WMA] associated with gross RV dilatation, findings that suggest global depress RV function and invasive hemodynamic monitoring identifying RVI by a combination of findings that suggest dysfunction of the RV [low cardiac output [CO] and a disproportionate elevation of the right atrial pressure[PWP]]]. The diagnosis for shock was made if all the following criteria were met: SSP persistently ≤ 90 mm Hg or vasopressors required to maintain SSP > 90 mm Hg; very low CO [cardiac index [CI] < 2.1L.min/m²]; evidence of end organ hypoperfusion.

Clinical classification and the TIMI-RS

We risk-stratified RVI patients into 3-subsets based on clinical features and echocardiographic findings upon admission, that were further supported with hemodynamic findings as follows: Class A, comprised patients without evidence of systemic hypotension [SSP ≥ 100 mm Hg] or RVF [by clinical features, echocardiographic and/or hemodynamic findings]; Class B, were those with persistent systemic hypotension [SSP < 100 mm Hg] or RVF, but without other clinical features of shock; and Class C, were those in shock. TIMI-RS was calculated according to previously published criteria. We tested retrospectively 2-schemes; TIMI-RS and the CC for 425 RVI patients; prospectively between January 2003 and July 2007 among 622 consecutive patients with STEMI [anterior: 277; inferior: 247; RVI: 98].

Reperfusion procedures

For RVI patients, considered for thrombolytic therapy [TT] or percutaneous coronary intervention [PCI]; TT was administered to all eligible RVI patients; inclusion criteria for classes A and B were defined before admission as follows: symptoms of MI lasting less than 6h; age ≤ 75y; and absence of other well known accepted contraindications. TT was performed with either streptokinase or recombinant tissue plasminogen activator, preceded by a heparin bolus. Adjunctive treatment consisted of an intravenous heparin dose that was adjusted to keep the TPTa between 60-80s and aspirin. For PCI the same inclusion criteria for MI symptoms duration/age were applied; stent deployment was performed according to standard techniques and follow by the use of standard anti-platelet therapy. Stents and glycoprotein IIb/IIIa platelet inhibitors became standard therapy in eligible patients in 1995 and 1996, respectively. Clopidogrel was used in stented patients [loading dose 300 mg] and continued [75 mg] for ≥ 6 months in patients treated with bare metal stents and for 6 to 12 months in patients treated with drug-eluting stents. For those RVI patients in shock considerer candidates for PCI: age < 75y and shock within < 24 hours of evolution. Conservative therapy for A and B classes was follow due to: symptoms of MI lasting more than 6h and/or to delay (> 6h) in patients arrivals for seeking for medical attention; for class C patients for those with age > 75y or to those who develop shock > 40h of the onset of MI.

Primary end point was in-hospital cardiac death at 20-30 days. Follow-up information after hospital discharge was obtained from the hospital records from the Instituto Nacional de Cardiología “Ignacio Chávez” registration database that is updated at each patient visit and upon patient death.

Statistical analysis

Continuous data are expressed as mean ± SD. Student t test, 1-way ANOVA [Bonferroni’s test for multiple comparison], χ², or Fisher exact
test was used as appropriate. Univariate analysis based on the logistic regression model was used to examine the relation between selected demographic, medical history, clinical examination and ECG/ECHO data to add points in attempt to building-up the TIMI-RS-RVI model. Results are expressed as odds ratios 95% confidence intervals [CI]. At completion of the univariate analysis, any variable whose univariate test had a value of $p < 0.25$ was considered a candidate for the multivariate model. Inclusions into the multivariate logistic regression model was determined by the number of variables associated with a significance level of $p < 0.05$ in a stepwise elimination process were retained in the final model. After the predictor variables in the regression model had been finalized, the prognostic discriminatory capacity for the TIMI-RS-RVI was expressed as the c statistic, representing the area under the receiver operating characteristic [ROC] curve for prediction of in-hospital death. For the survival analysis we used the time of the initial ECHO/RHC as an index for determining survival. The Kaplan-Meier method was used to estimate overall survival distribution [Log-rank test]. Analyses were performed by using of SPSS-13 and STATA-9 software.

**Results**

The diagnosis of RVI was made by ECG and either echocardiographic, hemodynamic or coronary angiographic criteria in 100% and by three criteria [ECG, echocardiographic and hemodynamic] in 85% of the patients.

**Clinical and echocardiographic data**

The frequency in the baseline clinical characteristics used as predictors in the studied STEMI population and in the InTIME-II study are shown in Table I. Class A, B and C RVI-subgroups, had no differences in their baseline characteristics for both prospectively and retrospectively RVI studied patients. The echocardiographic data that give support to the CC for all studied RVI patients were documented as follows: no RV-dilatation, RV wall motion abnormalities [RVWMA] only for the inferior wall [IW = 100%] in addition to the free wall [FW = 14%], paradoxical ventricular septal motion [PVSM = 25%], tricuspid regurgitation [TR = 17%] with a normal LVEF (> 0.5) in all class A patients. For class B and C patients in all of them was demonstrated RV-dilatation and RVWMA not only confined to the IW. TR and PVSM were present in 78 and 70% [respectively] and in 29% of Class C patients an abnormal LVEF was documented.

**Hemodynamic data**

For the total RVI patients, invasive hemodynamic evaluation and coronary angiograms were performed in 81% and 76%, respectively. The hemodynamic data according to RVI Class was documented as follows: for class A patients \([n = 235]\) : mean right atrial pressure \([\text{mRAP} = 4.6 \pm 2.1 \text{ mm Hg}]\), systolic pulmonary artery pressure \([\text{sPAP} = 16.8 \pm 4.4 \text{ mm Hg}]\), diastolic \([\text{dPAP} = 9.7 \pm 3.7 \text{ mm Hg}]\), mean PWP \([8.6 \pm 3.1 \text{ mm Hg}]\), CI = 3.4 ± 0.71 L.min/m², mean systemic arterial pressure \([\text{mSAP} = 108.8 \pm 8.7 \text{ mm Hg}]\] were within normal hemodynamic limits. RAP/PWP ratio

| Table I. Baseline clinical characteristics used as predictors in the InTIME-II TIMI-RS and in the studied STEMI population. |
|---|---|---|---|
| Variables | InTIME-II | STEMI | RVI |
| Age ≥ 75y | 13.7 | 11.1 | 12.8 | 10.8 |
| Age 64 –74y | 41.9 | 32.7 | 35.2 | 37.5 |
| Diabetes | 13.9 | 28.8 | 24.1 | 30.6 |
| Hypertension and angina | 30.4 | 44.3 | 44.1 | 44.4 |
| Clinical findings | | | | |
| SSP < 100 mm Hg | 2.6 | 15.8 | 33.7 | 28.0 |
| Heart rate > 100 min | 7.7 | 14.4 | 12.6 | 13.8 |
| Killip Class II-IV | 12.6 | 29.1 | 31.7 | 26.0 |
| Weight < 67 kg | 19.2 | 22.2 | 12.2 | 11.5 |
| Presentation | | | | |
| Anterior ST elevation/LBBB | 42.7 | 52.8 | 0.0 | 0.0 |
| Time to therapy > 4h | 24.3 | 66.6 | 55.3 | 50.9 |

LBBB: left bundle branch block; SSP: systemic systolic pressure.
was found ≥ 0.8 in 2% of the RVI class A patients. Hemodynamic measurements after volume loading with normal saline solution [increments of 200ml, range 200 to 1,400] until the mean PWP was 15-19 mm Hg were perform in 32% class A patients, where the volume loading maneuver ratify the assigned RVI classification at arrival for this patients. For class B patients [n = 119] an elevated mRAP [12.9 ± 3.6 mm Hg], decreased CI [2.4 ± 0.21 L.min/m²] and mSAP [78.7 ± 12.3 mm Hg] were documented with an increase in the RAP/PWP ratio [96% of the patients]; p < 0.01 between A and B RVI classes. For class C patients [n = 69] an elevated mRAP [21.4 ± 5.15 mm Hg], sPAP [36.8 ± 9.3 mm Hg], dPAP [22.2 ± 3.9 mm Hg] and mPWP [19.9 ± 9.2 mm Hg], decreased CI [1.67 ± 0.5 L.min/m²] and mSAP [62.7 ± 9.5 mm Hg] were documented with an increase in the RAP/PWP ratio [72% of the patients]; p < 0.01 between B and C RVI classes.

**Angiographic results**
The right coronary artery [RCA] was the infarct related artery in 93% of the studied RVI patients [n = 397]. The 3-CC-RVI subgroups had no differences in the number of diseased vessels. Complete RCA obstruction was found in 210 patients and 53% of these had coronary collateral circulation. Significant differences in coronary collaterals were found between A, B and C patients [52.4, 48.3 and 8.2%; respectively; p < 0.04]. The combination of RCA and significant left anterior descending [LAD > 50% stenosis] disease was more commonly found in C patients [58.7%], which was significantly different from A [19.6%] and B [29.9%] patients [p < 0.03].

**Reperfusion results**
From the 523 RVI patients, 359 [68%] were considered eligible for thrombolytic therapy [TT] or percutaneous coronary intervention [PCI]. TT was performed with either streptokinase [32%] or recombinant tissue plasminogen activator [68%]. According to the current concepts of success by TT reperfusion⁹-¹¹ or by PCI,¹² this was achieved as a mean average in 73.6% of the patients. Mortality in these patients according to their class was as follows. When comparing with/without reperfusion for A, B and C classes, mortality was reduced when TT or PCI was administered [7.7% versus 3.5%, p < 0.05; 26% versus 11%, p < 0.01; 89% versus 50%, p < 0.01, respectively].

**Morbidity and in-hospital mortality**
Most class A patients had an uneventful clinical course [60.5%]. The remaining patients developed one or two of the following complications: sustained hypotension lasting less than 12h [18.7%] and cardiogenic shock. For class B patients, reversible RV dysfunction [defined as normalization of the SSP without volume or inotropes and/or improvement or normalization of RVWMA and RV dilatation on ECHO] was documented in 82% patients, all of whom survived. Most of the remainder patients developed cardiogenic shock. For class C patients, it was observed a significant major incidence of third degree AV block [pacemaker requirement] when compared to A and B classes [57 versus 5 and 18%, respectively; p < 0.04].

**The TIMI-RS analysis**
Application of TIMI-RS retrospectively for RVI patients did not reveal an association between outcome and TIMI-RS, nor did it predict short-term mortality [Fig. 1]. The lack of association of mortality with TIMI-RS was observed in both the low [≤ 4 points] or higher score [≥ 5] groups. When the TIMI-RS was tested prospectively in 98 RVI patients, the score did not reveal a significant graded increase in mortality with rising score [Fig. 2]. TIMI-RS showed a direct relationship with mortality in the 524 STEMI patients without-RVI that were evaluated prospectively [Fig. 2, range 3%-28.9%; p < 0.001 for trend].

![In-hospital mortality by TIMI risk score analysis in RVI patients](image-url)
To establish whether TIMI-RS should incorporate clinical points for right-sided electrocardiographic changes the following variables were tested: preinfarction angina, syncope, elevated neck venous pressure, Kussmaul’s sign, right-sided S₃/S₄, ST-segment elevation [<1 mm], new Q wave in V₃R/V₄R, RBBB, bradycardia, third degree AV block, ventricular arrhythmias, clear lung fields on chest X-rays and echocardiographic evidence of RV dilatation. This process yielded the following independent variables: age $\geq 75$y, age 64-74y, SSP < 100 mm Hg, heart rate > 100 min, Killip class II-IV, preinfarction angina, ST-segment elevation $\geq 1$ mm in V₄R, syncope, third degree AV block and elevated neck venous pressure [Table II]. When submitted to stepwise logistic regression: age 64-74y, age $\geq 75$y, SSP < 100 mm Hg, heart rate > 100 min, Killip class II-IV remained as a significant predictors of mortality in the multivariate model [Table III]. For the derived model to predict the probability for mortality [TIMI-RS-RVI] a c statistic = 0.9 was found, value that indicate a model with an excellent discrimination, but with a “very poor calibration”.

**Table II.** Results of univariate predictors for clinical and right-sided precordial electrocardiographic variables included for testing points in the TIMI-RS-RVI [n = 425].

<table>
<thead>
<tr>
<th>Historical</th>
<th>OR [95% CI]</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age $\geq$ 75 years</td>
<td>4.0 [1.8–9.0]</td>
<td>$&lt; 0.001$</td>
</tr>
<tr>
<td>Age 64 -74 years</td>
<td>4.4 [2.2–9.2]</td>
<td>$&lt; 0.001$</td>
</tr>
<tr>
<td>Diabetes+Hypertension+Angina</td>
<td>1.8 [0.6-2.2]</td>
<td>0.788</td>
</tr>
<tr>
<td>Prior angina</td>
<td>5.5 [2.2–14.8]</td>
<td>$&lt; 0.000$</td>
</tr>
<tr>
<td>Syncope</td>
<td>4.3 [2.0–10.6]</td>
<td>0.001</td>
</tr>
<tr>
<td>Examination/ECG/ECHO</td>
<td>3.7 [1.9-8.4]</td>
<td>$&lt; 0.000$</td>
</tr>
<tr>
<td>SSP &lt; 100 mm Hg</td>
<td>2.2 [1.2–3.0]</td>
<td>$&lt; 0.000$</td>
</tr>
<tr>
<td>Killip Class II-IV</td>
<td>3.2 [2.5–4.1]</td>
<td>$&lt; 0.000$</td>
</tr>
<tr>
<td>Weight &lt; 67 Kg</td>
<td>1.4 [0.4–2.7]</td>
<td>0.962</td>
</tr>
<tr>
<td>Neck vein distention</td>
<td>2.3 [5.0–28.6]</td>
<td>$&lt; 0.000$</td>
</tr>
<tr>
<td>Without pulmonary congestion</td>
<td>3.8 [0.8–3.7]</td>
<td>0.085</td>
</tr>
<tr>
<td>Bradycardia</td>
<td>2.0 [0.7-4]</td>
<td>0.40</td>
</tr>
<tr>
<td>ST elevation/Q wave–V4R</td>
<td>3.7 [1.5–9.3]</td>
<td>0.02</td>
</tr>
<tr>
<td>Third degree AV Block</td>
<td>3.8 [1.9–7.6]</td>
<td>0.001</td>
</tr>
<tr>
<td>RV dilation on ECHO</td>
<td>1.4 [0.6–2.9]</td>
<td>0.475</td>
</tr>
<tr>
<td>Presentation</td>
<td>1.4 [0.6–2.4]</td>
<td>0.578</td>
</tr>
</tbody>
</table>

RV: right ventricular, other abbreviations as in Table I.

**Table III.** Odds Ratios for predicting in-hospital mortality obtained from the multivariate logistic regression analysis.

<table>
<thead>
<tr>
<th>Variable</th>
<th>B</th>
<th>SE</th>
<th>Wald</th>
<th>df</th>
<th>OR</th>
<th>95%CI</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age 64-74y</td>
<td>1.473</td>
<td>.578</td>
<td>6.487</td>
<td>1</td>
<td>4.4</td>
<td>1.4-14.5</td>
<td>$&lt; 0.01$</td>
</tr>
<tr>
<td>Age 75-80y</td>
<td>3.159</td>
<td>.666</td>
<td>22.503</td>
<td>1</td>
<td>23.9</td>
<td>6.6-82.4</td>
<td>$&lt; 0.001$</td>
</tr>
<tr>
<td>SSP &lt; 100 [mm Hg]</td>
<td>1.906</td>
<td>.583</td>
<td>10.679</td>
<td>1</td>
<td>6.8</td>
<td>2.1-21.1</td>
<td>0.001</td>
</tr>
<tr>
<td>Killip Class II-IV</td>
<td>3.655</td>
<td>.568</td>
<td>41.366</td>
<td>1</td>
<td>37.5</td>
<td>12.7-117</td>
<td>$&lt; 0.01$</td>
</tr>
</tbody>
</table>

B: coefficient; SE: standard error; Wald: Walds test; df: degrees of freedom.
A progressive rise in mortality was seen: A: 5.4%, B: 16% and C: 50% [A versus B, p < 0.01; B versus C, p < 0.01]. The number of RVI patients alive at 8 years was for A class: 118, for B:60 and for C:10 patients. The significant differences between the RVI Classes that were observed in the short-term analysis were sustained at 8 years of follow-up [p < 0.001]. At 8 years, total mortality without intervention or failure in reperfusion in Class A patients was 35%, in B patients 55% and in C patients 90% [Fig. 4, solid lines]. When reperfusion therapy at admission was successful, total mortality at 8 years was lower than the previously mentioned for each RVI Classes [for A: 19%, for B: 32% and for C: 78%; p < 0.001, dashed lines].

Discussion

The studied RVI patient population where the CC and the TIMI-RS was tested is in accordance to cases described by other authors with STEMI for RV location, as defined by validated diagnostic criteria.12,13 Also our RVI patients were similar in regard to baseline demographics, treatment strategies and mortality to those described in the majority of previous studies.3,4,14,15 From 523 patients with RVI, 65% had silent hemodynamic RVI, 22% had RVF and 13% were in shock. Therefore, it should be emphasized that we covered all the “clinical spectrum of acute RVI” in the studied cohort. The TIMI-RS predictive value has been validated in a large, non-selected registry of STEMI patients.1,2 This observation was confirmed in our prospectively studied STEMI population with an anterior and inferior STEMI location. According to Gumina et al,5 their retrospective observations extended the utility of the TIMI–RS for RVI patients. Nonetheless, in the Gumina study5 the TIMI-RS helped mainly to stratify the risk in the minority of RVI patients with low scores, but failed to further stratify a large group of RVI patients with scores \( \geq 4 \). Also, when the mortality rate of RVI patients in their study is stratified by the TIMI-RS and compared with the mortality rate in patients with STEMI in the InTIME–II trial, large differences were observed.1,7 The analysis of both our whole RVI cohort [three-fold size of Gumina’s],5 or the prospective RVI cohort [similar in size of Gumina’s; 98 versus 102 patients, respectively]5 demonstrates a poor prognostic performance of the TIMI-RS. Accordingly, the lack of its applicability is particularly relevant in a cohort that comprises the wider clinical/hemodynamic spectrum for RVI. The divergent findings on the TIMI-RS for RVI patients could be due to the following explanations: 1- In TIME-II TIMI-RS was derived among patients with left ventricular STEMI [42.7% anterior, 56.9% inferior, others 0.4%]; therefore a small proportion of RVI patients where included for testing risk stratification, none were in shock and all were fibrinolytic-eligible patients.1 On the contrary, in our study all patients had RVI, no more than 68% underwent reperfusion thera-

Fig. 3. The Clinical Classification reveals a strong association between the short- and long-term risk of mortality for RVI patients.

Fig. 4. Significant differences between the 3-RVI Classes without intervention/failure [solid lines] or with success [dashed lines] of reperfusion therapy that were observed in the short-term were sustained at 8 years of follow-up.
py and 13% were in shock. Thus, differences in infarct location, area of myocardium at risk, time to therapy and hemodynamic status, could account for the different results derived from the TIMI-RS when applied to a non–selected population of RVI patients in clinical real life. 2- An arbitrary dichotomous cut-off value of SSP < 100 mm Hg which is used as a one of the major constituents in the TIMI score\textsuperscript{1} model–leave little room for further risk stratification of a population of RVI that covers the whole clinical spectrum and where the majority of RVI patients have a SSP ≥ 100 mm Hg [86%]. On the contrary, for our proposed classification, while using the same SSP-cut-off value it allows for the risk stratification of RVI patients in those without RVF, those with RVF or with cardiogenic shock.

Although, we found a low performance for the TIMI-RS in RVI patients we could not categorically denied the utility of this well proved tool on the basis of the following reasons [limitations]: 1.- it has been shown previously that the TIMI-RS has diminished performance in patients not treated with reperfusion therapy [31% of the studied population]. 2.- due to the mainly retrospective nature of the study we could also rise the question as to whether the TIMI-RS was always correctly determined and 3.- the information presented is coming from a single-cardiovascular center study.

Several investigators demonstrated that age, ST elevation in V4R, shock, ventricular arrhythmias and AV-block are associated with an increased risk of death for RVI patients.\textsuperscript{3,4,12,17} When we incorporate these variables for the construction of the TIMI-RS–RVI model in addition to echocardiographic abnormalities used for the diagnosis of RV dilation in the multivariate analysis; none except age [≥ 64y] and those hemodynamic conditions related to shock, remain as predictors of short-term mortality. The addition of these variables result in a TIMI-RS–RVI model with an excellent discrimination, but with a very poor calibration\textsuperscript{18} therefore, the derived model resulted in a measure of “non goodness-of-fit” and therefore could not be applied in clinical practice for RVI risk stratification. It is also seems the reason why, shock in this model failed to predict mortality when shock as part of the CC scheme did.

The long-term prognosis of RVI patients has been defined by several investigators.\textsuperscript{3,5,19,22} Our data reiterate that patients with RVI in the categories of RVF and cardiogenic shock have the high in-hospital morbidity and mortality.\textsuperscript{3,4,19,21,23–25} Accordingly, our information underscores the importance of making the correct stratification of RVI, with particular emphasis to consider at all times the “clinical spectrum of RVI” in its discussion.

The Global Use of Strategies to Open Occluded Coronary Arteries [GUSTO–I] risk model uses a graded scoring system for hemodynamics parameters that might be better for classification of patients at varying risk such as those with and without RVI.\textsuperscript{22} This hypothesis will need to be tested in RVI patients;\textsuperscript{7} in relation to, we should mention that we clearly separated RVI patients from those with only inferior-STEMI wall, and the classification for RVI patients was supported by hemodynamic findings in 80% RVI patients. In the present study, we report that long-term mortality continues to increase after the first year and is different for the 3-RVI Classes. The differences noted for long-term survival after RVI among the 3-Classes in the present study could be attributed to coronary artery disease progression, in part to the association of complete RCA obstruction significant LAD disease and a poorly developed collateral coronary circulation, conditions most frequently observed on admission for B and C RVI patients; and also according to our results, to the lack of indication and to the impact at admission of reperfusion success or failure.\textsuperscript{13,17,20}

The proposed Classification for RVI patients, which is based on the pathophysiology of RVI, is a simple, intuitive to most clinicians, non-invasive tool that can easy be applied at the bedside, has been supported by RV hemodynamic measurements that separate hemodynamically stable from unstable patients in the acute setting of RVI.

**Conclusions**

Although, the proposed Classification for short- and long-term survival is derived from retrospective and prospective data of a single cardiovascular center, our findings seems to support the use of a “Clinical Classification” on admission for risk stratification for acute RVI patients. The performance of the TIMI-RS in the acute setting of RVI should be investigated and reevaluated prospectively in a large cohort of RVI patients to ascertain its definitive role.
References


