High adherence to therapy and low cardiac mortality and morbidity in patients after acute coronary syndrome systematically managed by office-based cardiologists in Germany: 1-year outcomes of the ProAcor Study

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Abstract: We aimed to assess patient acceptance and effectiveness of a 12-month structured management program in patients after an acute coronary syndrome (ACS) event who were treated in a special setting of office-based cardiologists. The program comprised patient documentation with a specific tool (Bundesverband Niedergelassener Kardiologen [German Federation of Office-Based Cardiologists] cardiac pass with visit scheduling) shared by the hospital physician and the office-based cardiologist, the definition of individual treatment targets, and the systematic information of patients in order to optimize adherence to therapy. Participating centers (36 hospitals, 60 office-based cardiologists) included a total of 1,003 patients with ACS (ST-segment elevation myocardial infarction [STEMI] 44.3%, non-ST-segment elevation myocardial infarction [NSTEMI] 39.5%, unstable angina pectoris [UA] 15.2%, and unspecified 1.0%). During follow-up, treatment rates with cardiac medication remained high in all groups, with dual antiplatelet therapy in 91.0% at 3 months, 90.0% at 6 months, and 82.8% at 12 months, respectively. Twelve months after the inclusion, a total of 798 patients (79.6%) still participated in the program. Eighteen patients (1.8%) had died after discharge from hospital (6 in the STEMI, 12 in the NSTEMI group), while for 58 the status was unknown (5.8%). Based on a conservative approach that considered patients with unknown status as dead, 1-year mortality was 7.6%. Recurrent cardiac events were noted in 14.9% at 1 year, with an about equal distribution across STEMI and NSTEMI patients. In conclusion, patients’ acceptance of the ProAcor program as determined by adherence rates over time was high. Treatment rates of recommended medications used for patients with coronary heart disease were excellent. The 1-year mortality rate was comparatively low.

Keywords: myocardial infarction, patient management, patient education, feedback, patient-oriented outcomes, therapy adherence, compliance, mortality, quality of life

Introduction

Despite recent decreases in mortality rates in many countries, coronary heart disease (CHD) is still responsible for 1.8 million deaths in Europe, accounting for 20% of the total mortality. 1 ST-segment elevation myocardial infarction (STEMI), non-ST-segment elevation myocardial infarction (NSTEMI), and unstable angina pectoris (UA) form the group of acute coronary syndromes (ACS). Between 2005 and 2009,
ACS accounted for around 1,200 per 100,000 hospitalizations each year in Germany. During the hospital stay, around 7% of STEMI patients, 4% of NSTEMI patients, and 1% of UA patients die. Also after discharge, NSTEMI is associated with the highest mortality rate owing to a recurrent ischemic event: 6.2% of NSTEMI patients, 4.8% of STEMI patients, and 3.6% of UA patients do not survive within a 6-month period. After 5 years, these numbers increase to 22%, 19%, and 17%, respectively.

These epidemiological data show that it is crucial for patients after an ACS event to follow a stringent risk management regimen for prevention of secondary coronary events. To achieve this, the respective guidelines recommend lifelong treatment with drugs that repress thrombocyte function (aspirin and/or P2Y12 antagonists), decrease elevated low-density lipoprotein (LDL) cholesterol levels (statins), and lower elevated blood pressure (beta blockers and inhibitors of the renin–angiotensin–aldosterone [RAS] system). More than a decade ago, data from two registries (GRACE and CRUSADE) showed a clear association between adherence to treatments and survival.

From the patient’s perspective, as well as lifestyle changes, there is a long-term burden through the daily requirement to take multiple medications, although the effect of treatment is not apparent, as in the best case, coronary events do not occur. Conversely, adverse drug reactions may compromise the patient’s quality of life. Thus, independent of medication type, the adherence to treatment often is low, in particular if long-term therapy is necessary. Better translation of treatment guidelines into clinical practice with the aim of improving their acceptance by patients may substantially improve the effectiveness of current ACS treatment.

Among other measures, this may be achieved by a structured patient management approach, one of the aims of which is to improve cooperation between the acute clinic and the office-based cardiologists. Further, patients should be educated comprehensively about the implications of their chronic disease and receive regular feedback about the success of therapy. Smaller studies in Germany suggest that such approaches actively involving patients may help reach therapy targets.

In Germany, an existing program offered by the Bundesverband Niedergelassener Kardiologen [German Federation of Office-Based Cardiologists] (BNK) comprises patient documentation with a specific instrument (BNK cardiac pass with visit scheduling) done jointly by the hospital physician and the office-based cardiologists, the definition of treatment targets and the structured information of patients in order to optimize adherence to therapy. To gain additional evidence on this long-term ACS care program that engages patients, the ProAcor study was implemented. It observes patients’ adherence to and the effectiveness associated with this program.

The outcomes of this program are here reported as with focus about patients’ adherence to medication, treatment patterns, and outcomes under real-life conditions. The primary goal of the ProAcor study was the assessment of patients’ willingness to be guided through the structured outpatient care program. Further, it documented drug utilization patterns, patient satisfaction, and effectiveness measured by mortality and cardiac morbidity.

**Methods**

**Study sites and participants**

ProAcor is an open, nonrandomized, and observational prospective study that was set up to investigate patients with ACS treated under real-life conditions (outside clinical studies). A total of 36 clinics and hospitals and 60 resident cardiologists from all German federal states participated in the ProAcor study. This study was approved by the Ethics Commission of the Bavarian Medical Chamber on November 8, 2011 (project number 11110); pertinent guidelines for data protection were respected. The study identifier in ClinTrials.gov is NCT01490645.

Hospital and office-based cardiologists were entitled to participate if they treated a significant number of ACS patients and had sufficient personnel and trial experience for participation in the study. They also needed to have access to the web-based “Electronic Quality Management System (eQM)” of BNK Service GmbH.

Patients were eligible for documentation (inclusion criteria) if they were at least 18 years old; had been hospitalized owing to a STEMI, NSTEMI, or UA event; and had provided written informed consent for study participation within 7 days of the ACS index event.

Patients could not be documented if one or more of the following criteria were met (exclusion criteria): 1) UA, STEMI, and NSTEMI precipitated by or as a complication of surgery, trauma, or gastrointestinal bleeding or post percutaneous coronary intervention (PCI). 2) UA, STEMI, and NSTEMI occurring in patients already hospitalized for other reasons. 3) Presence of any condition/circumstance that in the opinion of the investigator could significantly limit the complete follow-up of the patient (eg, tourist, nonnative speaker or one who does not understand the local language, known diagnosed psychiatric illness, active cancer disease).
4) Patient not intended to be treated according to current guidelines, including patients participating in a clinical trial.

**Study flow and components**

The study flow including the study components is shown in Figure 1. In participating hospitals, the treating physician approached consecutive suitable patients, informed them about the goals and requirement of the study, and obtained written consent. The participation in the study did not change the way the patient was treated.

During the hospital phase of the study, patients obtained the Cardio Pass (BNK Herzpass®, Munich, Germany) of the German Federation of Office-based Cardiologists. This booklet is used for the shared documentation of the patient’s clinical history by the hospital physician and the office-based cardiologist with the aim of facilitating transfer of relevant information and agreement on common goals of secondary prevention. The Cardio Pass also includes a visit planner. Further, the patients receive information brochures and cards about their disease and recommended lifestyle changes to prevent further cardiac events after hospital stay.

After discharge from hospital, patients visited their office-based cardiologists for examinations and feedback about their state of health at intervals of 3, 6, and 12 months. At each of these three visits, the patient’s individual treatment goals (in terms of antiplatelet and other cardiac drug use, blood pressure, lipid and blood glucose values) and recommended lifestyle changes were assessed and discussed between the patient and the treating (office-based) cardiologist. At these visits, the Cardio Pass was updated with therapy goals (which could be modified during follow-up, if needed) and medication changes. To remind the patient about the lifestyle and treatment goals, information cards were handed out (after 9 months sent via mail) every 3 months during the program duration.

**Patient satisfaction**

To evaluate patient satisfaction with the ProAcor patient management program, a proprietary questionnaire was developed that stated the questions displayed in Table 1.

**Patients’ medication adherence**

Physicians rated the adherence to drug therapy on the basis of their own subjective assessment (without specifically asking the patients) after 12 months. They had the options “always (7 days per week)”, “mostly (4–6 days per week)”, “rarely (1–3 days per week)”, and “never (0 days per week)”.

**Data collection and analysis**

At all visits, data were collected from the patient charts into the Electronic Quality Management System (eQM), a central database. Collected parameters are summarized in Table 2. Data were documented in a pseudonymized way (i.e., only the treating physicians knew their patients’ identity). Measures for quality assurance comprised plausibility checks during data entry, statistical validation, and queries to centers if information was missing or ambiguous.

A sample size of 1,000 patients was originally planned to estimate the patient acceptance rates of the program (primary end point) with a precision of 5%.

An exploratory–descriptive statistical analysis covering parameters (qualitative, quantitative, and text fields including derived and coded variables) from the electronic case report forms (CRFs) was performed. Binary, categorical, and ordinal parameters were summarized by means of absolute and percentage numbers within the various categories, including “missing data” as a valid category at the baseline visit. Numerical data were summarized by means of standard statistics (ie, number of available data, number of missing data, mean, standard deviation, minimum, median, maximum, and lower and upper quartile). Where useful, the summary statistics were presented by visit. Calculations were performed with nQuery Advisor 7.0.

**Results**

**Patient disposition and flow**

A total of 1,003 patients were included and at least one visit documented (full analysis set). In 11 patients, the type of ACS was not specified. Eight hundred eighty-seven patients had information about the ACS type and had at least one
follow-up visit (evaluable patients set). One year after the inclusion, a total of 798 patients (90.0%) still participated in the program, while 89 patients (10.0%) did not. Dropout rates were somewhat higher in the STEMI group (11.1%) and the UA group (12.0%), respectively, compared to the NSTEMI group (8.1%). Mean participation time in the program was 11.4 ± 2.0 months overall.

Reasons for premature withdrawal from the program in the 89 patients were change of their medical doctor in 10 cases, withdrawal of informed consent in 14 cases, patient lost to follow-up in 57 cases, death of the patient in 3 cases, and patient relocation in 5 cases.

Demographics, characteristics, and concomitant diseases at baseline
Of the evaluable patients, male patients prevailed (73.6%; Table 3 top). The rate of males was higher in the STEMI and UA groups compared to the NSTEMI group. Mean age of patients was 61.7±11.6 years overall; in the STEMI group it was somewhat lower (58.9±11.2 years) than in the NSTEMI group (64.0±11.6 years) and in the UA group (64.3±10.9 years). Mean body mass index was 28.4±4.7 kg/m², with no relevant differences between groups.

The most frequently noted concomitant diseases were arterial hypertension (72.6%), hyperlipoproteinemia (59.2%), diabetes mellitus (22.2%), and renal insufficiency (7.2%) (Table 3 center). Current smoking was noted in 29.1% of patients, previous smoking in 27.6%. Valve disorders were reported in 6.8% and heart rhythm disorders (most frequently atrial fibrillation) in 7.4%. Patients in the NSTEMI group tended to have a higher prevalence of concomitant diseases and cardiovascular risk factors compared to patients in the STEMI and UA groups, respectively.

A prior cardiac event was noted in 15.3% overall, with higher frequency in the NSTEMI and UA groups compared to the STEMI group. Such events had led to earlier percutaneous transluminal coronary angioplasty (PTCA) in 12.1%, stent implantation in 11.1%, and cardiac bypass surgery in 1.9%, respectively (Table 3 bottom).

Index cardiac event and antiplatelet therapy
The index cardiac event was STEMI in 407 evaluable patients (of 444 total), NSTEMI in 347 patients (of 396 total), and UA in 133 patients (of 152 total).

Overall, for 750 patients (84.6%) the index event was their first manifestation of CHD. By subgroup, this was the case...
High adherence to therapy and low cardiac mortality and morbidity

for 370 patients in the STEMI group (90.9%), 289 patients in the NSTEMI group (83.3%), and 91 patients in the UA group (68.4%), respectively.

In the context of the ACS index event, 828 patients (93.3%) received a PTCA, and 801 patients received a stent. The rates of PTCA and stents, respectively, were higher in the STEMI group compared to the NSTEMI and UA groups, respectively.

After the index event, patients were discharged from the acute hospital after 6.8±8.4 days (median 6.0). Patients in the STEMI group stayed longer in the hospital (8.2±11.0, median 6.0) compared to patients in the NSTEMI group (6.1±5.2, median 5.0) and the UA group (4.0±3.5, median 3.0), respectively.

Antiplatelets and other cardiac medication

For the treatment of the acute ACS index event, almost all patients (n=858, 96.7%) received at least one P2Y12 inhibitor or acetylsalicylic acid (ASA), (Figure 2). The great majority (819 patients, 92.3%) received dual antiplatelet therapy (DAPT). DAPT treatment rates were higher in STEMI patients (395 patients, 96.1%) compared to NSTEMI patients (314 patients, 90.5%) and UA patients (110 patients, 82.7%). At the initiation of P2Y12 inhibitor therapy, the respective drug was provided as a loading dose in 406 patients (45.8%). Treatment rates with antiplatelet drugs remained high during the study follow-up. Of the evaluable patients at each visit, 812 patients (93.8%) after 3 months, 772 patients (92.7%) after 6 months, and 682 patients after 12 months (85.5%) received ASA/any of the P2Y12 inhibitors as monotherapy. DAPT was reported in 788 patients (91.0%) at 3 months, 750 patients (90.0%) at 6 months, and 661 patients (82.8%) at 12 months, respectively. For all time points, rates of antiplatelet drugs (any or DAPT) were higher in the STEMI group compared to the NSTEMI and UA groups.

Almost all patients received some cardiac medication after the index event. Drug classes most frequently

### Table 2: Collected patient parameters at baseline, follow-ups, and at the final examination

<table>
<thead>
<tr>
<th>Visit</th>
<th>Baseline*</th>
<th>Follow-Up</th>
<th>Final</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time (months)</td>
<td>0</td>
<td>Q1</td>
<td>Q2</td>
</tr>
<tr>
<td>Date of visit</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Informed Consent</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inclusion/exclusion criteria</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distribution of information brochure/cards</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hand out/update of BNK Herzpass®</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Arrange a follow-up visit at the BNK cardiologist (optional)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fax for the index documentation</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Last echocardiography findings</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Last vital values (blood pressure, heart rate, dyspnea)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Last laboratory values (troponin, potassium, creatinine, glucose, total/HDL/LDL cholesterol, lipoproteins, triglycerides, HbA1c)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Demographic data (date of birth, sex)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient history including CV risk assessment (smoking status, hyperlipoproteinemia, diabetes, hypertension, CHD family history)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date of ACS diagnosis and type of ACS</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACS index event-related procedures (interventions and medications)</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concomitant medication (identical/different from row above)</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Concomitant diseases</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Physical/cardiological routine examination</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>New cardiac events*</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Assessment of individual treatment goals/lifestyle changes</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Distribution of patient information cards*</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

**Notes:** Procedures performed at the hospital. *Source data for this information was hospital discharge letter (index event). †Any new episode of the ACS, stroke, CV procedure, hospitalization or death. ‡Patient information cards were handed out intuitively according to patients’ situation (eg, only current smoker should receive information on smoking cessation). §Patient information cards were sent out by the investigator, or BNK Service GmbH if desired by the investigator.

**Abbreviations:** BNK, Bundesverband Niedergelassener Kardiologen; HDL, High-density lipoprotein; LDL, Low-density lipoprotein; CV, Cardiovascular; CHD, coronary heart disease; ACS, acute coronary syndrome.
prescribed – besides platelet inhibitors in 858 patients (96.7%) – were beta blockers in 809 patients (91.2%), CSE inhibitors (statins) in 774 patients (87.3%), and ACE inhibitors in 611 patients (68.9%). Prescription rates of these drug classes are presented by ACS type in Figure 3.

During follow-up, treatment rates with cardiac medication remained high in all groups (at least one drug in 857 patients [99.0%] at 3 months, in 829 patients [99.5%] at 6 months, and in 795 patients [99.6%] at 12 months).

At all three time points, patients in the STEMI group received cardiac drugs more often than did patients in the NSTEMI and UA groups, respectively.

**Adherence to therapy**

Physicians rated the adherence of their patients after 12 months in 793 patients (100%). According to their (subjective) assessment, 698 patients (88.0%) were always adherent, 90 patients (11.3%) mostly adherent, 4 patients (0.5%) rarely

### Table 3 Demographic characteristics of patients, comorbidities, and interventions after the index ACS event

<table>
<thead>
<tr>
<th></th>
<th>STEMI (n=407)</th>
<th>NSTEMI (n=347)</th>
<th>UA (n=133)</th>
<th>Total (n=887)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Demographics</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years ± SD)</td>
<td>58.9±11.2</td>
<td>64.0±12.0</td>
<td>64.3±10.9</td>
<td>61.7±11.6</td>
</tr>
<tr>
<td>Sex, males</td>
<td>312 (76.7)</td>
<td>237 (68.3)</td>
<td>104 (78.2)</td>
<td>653 (73.6)</td>
</tr>
<tr>
<td><strong>Comorbidities</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arterial hypertension</td>
<td>271 (66.6)</td>
<td>273 (78.7)</td>
<td>100 (75.2)</td>
<td>644 (72.6)</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>78 (19.2)</td>
<td>78 (22.5)</td>
<td>41 (30.8)</td>
<td>197 (22.2)</td>
</tr>
<tr>
<td>Hyperlipoproteinaemia</td>
<td>235 (57.7)</td>
<td>206 (59.4)</td>
<td>84 (63.2)</td>
<td>525 (59.2)</td>
</tr>
<tr>
<td>Peripheral arterial occlusive disease</td>
<td>15 (3.7)</td>
<td>23 (6.6)</td>
<td>10 (7.5)</td>
<td>48 (5.4)</td>
</tr>
<tr>
<td>Renal insufficiency</td>
<td>22 (5.4)</td>
<td>34 (9.8)</td>
<td>8 (6.0)</td>
<td>64 (7.2)</td>
</tr>
<tr>
<td>Family history of CHD</td>
<td>161 (39.6)</td>
<td>106 (30.5)</td>
<td>43 (32.3)</td>
<td>310 (34.9)</td>
</tr>
<tr>
<td>Prior stroke</td>
<td>9 (2.2)</td>
<td>14 (4.0)</td>
<td>7 (5.3)</td>
<td>30 (3.4)</td>
</tr>
<tr>
<td>Prior TIA/PRIND</td>
<td>1 (0.2)</td>
<td>5 (1.4)</td>
<td>3 (2.3)</td>
<td>9 (1.0)</td>
</tr>
<tr>
<td>Prior cardiac event</td>
<td>37 (9.1)</td>
<td>57 (16.4)</td>
<td>42 (31.6)</td>
<td>136 (15.3)</td>
</tr>
<tr>
<td><strong>Interventions in hospital</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTCA</td>
<td>398 (97.8)</td>
<td>318 (91.6)</td>
<td>112 (84.2)</td>
<td>828 (93.3)</td>
</tr>
<tr>
<td>Stent</td>
<td>392 (98.5)</td>
<td>305 (95.9)</td>
<td>104 (92.9)</td>
<td>801 (90.3)</td>
</tr>
<tr>
<td>Drug eluting stent</td>
<td>265 (67.7)</td>
<td>221 (72.5)</td>
<td>70 (67.3)</td>
<td>556 (62.7)</td>
</tr>
<tr>
<td>Bare metal stent</td>
<td>70 (17.9)</td>
<td>59 (19.3)</td>
<td>24 (23.1)</td>
<td>153 (17.2)</td>
</tr>
<tr>
<td>Coronary artery bypass surgery</td>
<td>3 (0.7)</td>
<td>10 (2.9)</td>
<td>4 (3.0)</td>
<td>17 (1.9)</td>
</tr>
</tbody>
</table>

**Notes:** Values are n (%), if not specified otherwise.  
**Abbreviations:** STEMI, ST-segment elevation myocardial infarction; NSTEMI, non-ST-segment elevation myocardial infarction; UA, unstable angina; CHD, coronary heart disease; PTCA, percutaneous transluminal coronary angioplasty; SD, standard deviation; TIA, transient ischemic attack; PRIND, prolonged ischemic neurological deficit.

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**Figure 2** Antiplatelet therapy.  
**Notes:** Values show the prescription rates (%) of the named drug classes or drugs in the total cohort, and in the STEMI, NSTEMI and UA subgroups.  
**Abbreviations:** DAPT, dual antiplatelet therapy; STEMI, ST-segment elevation myocardial infarction; NSTEMI, non-ST-segment elevation myocardial infarction; UA, unstable angina.
High adherence to therapy and low cardiac mortality and morbidity

Figure 3 Cardiac medication other than antplatelets during the course of the observation.

Abbreviations: STEMI, ST-segment elevation myocardial infarction; NSTEMI, non-ST-segment elevation myocardial infarction; UA, unstable angina; RAAS, renin angiotensin aldosterone system; ACE-I, angiotensin converting enzyme inhibitor; ARB, angiotensin receptor blocker.
adherent, and 1 patient (0.1%) never adherent. There were no relevant differences in therapy adherence among patients from the STEMI, NSTEMI, or UA groups, respectively.

Patient satisfaction
Patients were highly satisfied with the information about the risk factors for their disease and about the way to treat it by a change of lifestyle (Table 1). Of the patients, 96.3% felt sufficiently informed on the reasons for their disease, and 94.6% felt sufficiently informed on the possibility of treating their disease via change of lifestyle.

Mortality and clinical events during follow-up
Including all cases for which a definite survival status was not reported, mortality during the 12-month observation period was 7.6% at most (18 confirmed deaths, in addition to 58 patients lost to follow-up and thus of unclear vital status).

After the index ACS event, new cardiac events occurred in 128 patients (14.4%, Figure 4) during the full observational period. Seventy-eight patients (9.0%) had the new event within the first 3 months, 30 patients (3.6%) between 3 and 6 months, and 35 patients (4.4%) between 6 and 12 months. The rate of new cardiac events was higher in the STEMI group (62 patients, 15.2%) than in the NSTEMI group (51 patients, 14.7%) and the UA group (15 patients, 11.3%), respectively.

Furthermore, the majority of reported events were angina pectoris (86 patients, 67.2%), STEMI (20 patients, 15.6%), NSTEMI (17 patients, 13.3%), or a combination of these events (5 patients, 3.9%). Eighteen of the 20 STEMI events occurred in patients in the STEMI group.

Discussion
The ProAcor study, reporting on almost 1,000 patients, with ACS treated first in 36 hospitals and later by 60 office-based cardiologists, assessed patients’ acceptance of a structured monitored program along with outcomes over 1 year. The adherence to the program was high.

In the first year after the index event, ACS patients have a high risk of recurrent cardiac events; thus, secondary prevention measures are crucial to avoid cardiovascular mortality and morbidity.6,17 It is known, however, that many patients have low therapy adherence and readiness for lifestyle changes.18 Non-adherence to medical plans is a pivotal public health problem at every level of the population, particularly in older adults.19

In the experimental setting of randomized controlled trials (RCT), patient conditions are strictly controlled, leading to an exclusion of patients with comorbidities and concomitant medication. In contrast, subjects enrolled for observational studies such as ProAcor are typical, unselected patients, with a substantial proportion of comorbidities, and, therefore, additional medications. This mirrors real-world conditions.
of treatment administration and thereby complements the results of RCTs. Patient selection, medication, and time of follow-up in the ProAcor study were similar to other registry studies with ACS patients such as the Austrian APCI,20 the Italian EYESHOT,21 the international EPICOR,22 the French FAST-MI2010,23 or Swedish SCAAR.24

Adherence and patient satisfaction
Many different approaches, including educational, behavioral, and affective categories, have been tested to improve patient compliance with and adherence to medical interventions. Most interventions associated with adherence improvements are not associated with improvements in other outcomes; indeed, in the last few decades, the degree of nonadherence has remained unchanged.19 In an older meta-analysis by Roter et al,25 no single strategy or programmatic focus showed any clear advantage compared with any other, and comprehensive interventions combining cognitive, behavioral, and affective components were more effective than single-focus interventions. A recent Cochrane review by Nieuwlaat et al26 assessed 182 RCTs on the effects of interventions intended to enhance patient adherence to prescribed medications for medical conditions, on both medication adherence and clinical outcomes. They concluded that effects were inconsistent from study to study, and only a minority of the RCTs with the lowest risk of bias improved both adherence and clinical outcomes. Current methods of improving medication adherence for chronic health problems are mostly complex and not very effective, so the full benefits of treatment cannot be realized.26 However, earlier studies in ACS patients have shown that multifaceted patient-centered interventions could improve adherence to cardioprotective medications; in particular, respectful collaborative communication can contribute to medication adherence after ACS hospitalization.27,28

In order to increase therapy adherence and effectiveness in everyday clinical practice, essential features of the ProAcor study were the use of the Cardio Pass (Herzpass) containing information about patient history, medication and planning of visits, and patient education with cards and brochures about significance and measures for prevention of secondary cardiac events. On the basis of the high satisfaction patients expressed with regard to the information on the disease and the optional measures to treat it through lifestyle changes and medications, the program can be called successful.

Effectiveness
Treatment rates of recommended medications used for patients with CHD were excellent. Lack of compliance with treatment, eg, with antiplatelet agents, has been shown to be predictive of secondary events –such as stent thrombosis29– in the follow-up of ACS patients.

In fact, the observed mortality of 1.8% (confirmed) or 7.6% (most conservative approach), respectively, is lower than in several current ACS registries. In the German MONICA/KORA acute myocardial infarction registry, mortality was 3.2% in the first year after hospital discharge.30 For the Spanish DIOCLES study, in-hospital mortality was 4.1%, and in patients alive at discharge (follow-up available for 97.1%), 6-month mortality was 3.8%.31 This was already an improvement compared to the previous MASCARA32 registry study, which registered 10.5% deaths among patients after the same time period. Similarly, in the four French FAST-MI registry studies, mortality fell from 20% (1995) to 9.8% (2010) 1 year after hospital discharge.33 The CZECH-2 registry reported a 1-year mortality of 7.9%, the European MULTIPRAC registry of 2.4%, the Swiss SPUM-ACS of 4.6%, and the Swedish SCAAR registry of 6.9% (all data based on personal communication).

ProAcor differs from the other mentioned studies in that it observed a special management program that schedules several follow-up examinations of ACS patients after hospital discharge and focuses on measures for patient information and education. A recent systematic review by the Cochrane Collaboration of the effects of educational components of cardiac rehabilitation, apart from improvement of QoL ratings, did not find statistically significant reductions of hospitalizations or mortality.34 In an earlier analysis by the US Agency for Healthcare Research and Quality, a branch of the Food and Drug Administration (FDA), of randomized trials of secondary prevention programs in CHD, measures for patients after hospital discharge diminished the frequency of recurring hospitalization by 15% and the risk of myocardial infarctions within 12 months by 17%.35 Reduced mortality rates became statistically significant only in a more long-term perspective of 24 or 60 months (47% and 24%, respectively). Probably, patient education works synergistically with newer antiplatelet medication regimens and rigorous adherence to them; nonpharmacological and pharmacological treatments together may lead to a measurable diminishment of mortality within 12 months, as found in the ProAcor study.

Overall, the varying proportions of patients with STEMI/NSTEMI/UA and the completeness of follow-up make comparisons between studies difficult, but the low rate gives confidence that the patients in Germany in ProAcor do no worse than in comparable countries.
Limitations of the study
ProAcor, like all registry-type observational studies, has limitations, among them different types of bias. Managing physicians are a positive selection of motivated and well-trained cardiologists; thus, outcomes of the structured program may not mirror the “average” clinical practice. Only 50%–60% of patients were followed up by a cardiologist. Moreover, participating patients themselves were likely more motivated than the average patient population to know more about their disease, to change their lifestyle, and read the offered materials in tight time frames. Less interested patients with a lower training and adherence level may have led to less favorable outcomes. Thus, as in other ACS registries, allocation or channeling bias may confound the relation between treatment and outcomes. The patient satisfaction questionnaire was developed by study group members and has not yet been formally validated. Social and cultural levels of the participants were not documented. Assessment of compliance with or adherence to drug therapy relied on the subjective assessment of physicians, not on other pill count, check of filled prescriptions at the pharmacy, or patient interviews. Also, owing to its limited size and relatively short duration, the ProAcor study does not allow statements on the adherence of patients and effectiveness beyond one year.

Conclusion
In conclusion, patients’ acceptance of the ProAcor program, as determined by adherence rates over time, was high. Treatment rates of recommended medications used for patients with CHD were excellent. The 1-year mortality rate was low compared to that in other contemporary registries.

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