Aspirin Dosing: A Patient-Centric Trial Assessing Benefits and Long-term Effectiveness (ADAPTABLE) Study Protocol

Version 1.0
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I. Study Overview & Goals

IA. Study Rationale

Every year 720,000 Americans have a heart attack, and nearly 380,000 die of atherosclerotic cardiovascular disease (ASCVD).\(^1\) Many of the patients who survive develop heart failure, stroke, and/or other cardiovascular complications. As such, patients with ASCVD and their caregivers suffer from substantial symptomatic, emotional, and functional difficulties. These patients often experience chest pain, shortness of breath, and fatigue, which can lead to significant distress and worsening quality of life. Rates of mental health illness like depression are high among both these patients and their caregivers; rates of depression may approach 66% in post-myocardial infarction (MI) patients.\(^2\)-\(^8\) Coronary heart disease alone costs the United States $108.9 billion each year.\(^4\) This total includes the cost of health care services, medications, and lost productivity.\(^1\)

Aspirin is a mainstay therapy for patients with ASCVD. Introduced as a medicinal product more than 100 years ago, aspirin significantly reduces ischemic outcomes such as myocardial infarction and stroke in patients with previous cardiovascular events and/or atherosclerosis at a cost of less than a cent per day. However, despite dozens of clinical trials involving more than 200,000 patients, the optimal dose of aspirin—the dose that is most effective in reducing ischemic events in the setting of secondary prevention, balanced by the potential for adverse events such as gastrointestinal bleeding—has not been determined in direct comparative-effectiveness trials. Observational studies and indirect comparisons of different doses of aspirin have yielded conflicting results. Although most studies have found that lower-dose aspirin is associated with less bleeding, these studies have provided contradictory evidence regarding the comparative effectiveness of low vs higher-dose aspirin in reducing ischemic events. Additional evidence raises the possibility that patients with different underlying characteristics may benefit most from different doses of aspirin.

To identify the optimal dose of aspirin for secondary prevention in patients with ASCVD, we propose a pragmatic clinical trial in which 20,000 patients who are at high risk for ischemic events will be randomly assigned in a 1:1 ratio to receive an aspirin dose of 81 mg/day vs. 325 mg/day. Study participants will be enrolled over 24 months. Maximum follow-up will be 30 months. The primary endpoint is a composite of all-cause death, hospitalization for MI, or hospitalization for stroke. The primary safety endpoint is hospitalization for major bleeding with an associated blood product transfusion.

The ADAPTABLE trial study design is shown in the schematic below.
I.A.1. ADAPTABLE Study Design

**Study design**

Patients with known ASCVD (i.e., MI, OR cath ≥75% stenosis of ≥1 epicardial vessel OR PCI/CABG)

AND ≥1 Enrichment Factor*

Identified through EHR (computable phenotype) by CDRNs (PPRN pts. already part of a CDRN are eligible)

Pts. contacted with trial information and link to eConsent; Treatment assignment provided directly to patient

Exclusion Criteria
- Age < 18 yrs
- ASA allergy or contraindication (including pregnancy or nursing)
- Significant GI bleed within past 12 mos
- Significant bleeding disorder
- Requires warfarin or NOAC or Ticagrelor

ASA 81 mg QD  
ASA 325 mg QD

Electronic F/U Q 3-6 months; supplemented with EHR/CDM/claims data

Duration: Enrollment over 24 months; maximum I/u of 30 months

Primary Endpoint: Composite of all-cause mortality, nonfatal MI, nonfatal stroke

Primary Safety Endpoint: Major bleeding complications

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Enrollment and follow-up of study participants will be conducted using highly streamlined methods, with electronic health record (EHR) data organized according to the recently developed PCORnet Common Data Model (CDM) format and stored in a PCORnet DataMart,

*Enrichment factors
- Age > 65 years
- Creatinine > 1.5
- Diabetes (Type 1 or 2)
- 3-vessel coronary artery disease
- Cerebrovascular disease and/or peripheral artery disease,
- EF <50% by echo, cath, nuclear study
- Current smoker

Enrollment and follow-up of study participants will be conducted using highly streamlined methods, with electronic health record (EHR) data organized according to the recently developed PCORnet Common Data Model (CDM) format and stored in a PCORnet DataMart, complemented where possible by existing data sources (Medicare claims data) and patient reported outcomes. Additional information will be collected via streamlined forms to be completed by participants either by Internet if they are able to access the Internet or by the Call Center at the Duke Clinical Research Institute (DCRI). This project constitutes the initial randomized comparative-effectiveness trial conducted by the National Patient-Centered Clinical Research Network (PCORnet; [http://www.pcornet.org/](http://www.pcornet.org/)).

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* Please note: throughout this protocol, the term “Common Data Model” (CDM) is used to refer both to the format used for standardizing and organizing information, and also as shorthand for EHR data that are extracted and stored using the CDM format within the PCORnet DataMart.
This trial will incorporate several essential aspects of the new genre of patient-centered comparative effectiveness trials:

1. By using existing data sources to gather baseline characteristics and a combination of existing data and patient-reported outcomes during follow-up, the trial will answer this critical question at a relatively low cost.
2. An Internet portal will enable the trial to collect and monitor data and enable mutual learning by both patients and clinicians, capitalizing on the frequent use of the Internet by the American public and clinicians.
3. The trial will not have a placebo control, but instead will provide all patients with active treatment at different doses, with monitoring to balance benefit and risk.
4. Patient-reported outcomes will be collected.
5. The evolving PCORnet infrastructure will be used to streamline administrative aspects of the trial, including centralization of institutional review board (IRB) functions and contracts, electronic consent and use of EHR data standardized into the CDM format.

I.B. Study Aims

We have defined the following specific aims for this study:

- **Aim 1**: To compare the effectiveness of two daily doses of aspirin (81 mg and 325 mg) in reducing a composite endpoint of all-cause death, hospitalization for nonfatal MI, or hospitalization for nonfatal stroke in high-risk patients with a history of MI or documented atherosclerotic cardiovascular disease (ASCVD). Secondary endpoints will be the components of the composite primary endpoint as well as coronary revascularization procedures (percutaneous coronary intervention [PCI] or coronary artery bypass grafting [CABG] performed during study follow-up. The primary safety endpoint will be hospitalization for major bleeding complications with an associated blood product transfusion.

- **Aim 2**: To compare the effects of aspirin in selected subgroups of patients, including women vs men, older vs younger patients, racial minority patients vs white patients, patients with vs. without diabetes, and patients with vs. without chronic kidney disease (CKD)

- **Aim 3**: To develop, refine, and evaluate the infrastructure for PCORnet to conduct multiple comparative-effectiveness trials in the future. This aim will be accomplished with a "phased-in" approach (as previously described) that will allow for an initial testing of the PCORnet infrastructure followed by adjustments to the trial operational plan to most efficiently accomplish Aims 1 and 2. Also, during the first year of the trial, we will be carefully monitoring the recruitment and enrollment patterns within and across CDRN’s and will be providing regular feedback reports to each CDRN to promote consistency in the recruitment practices. Potential metrics to evaluate the success of ADAPTABLE are listed below and will be finalized with the PCORnet leadership in the context of other performance measures in development PCORnet-wide:
I.B.1. Comparison to DCRI Standard Metrics for Clinical Trials

- Time to IRB approval
- Time to contract approval
- Time to first site activation
- Time to first patient enrolled
- Recruitment rate
- Retention
- Withdrawn consents
- Drug discontinuation
- Lost to follow-up
- Missed study contacts
- Data quality

I.B.2. PCORnet as a Network

- Ability to support widespread screening, contact, enrollment, and follow-up of patients across the networks

I.B.3. CDRN Experience

- Administrative simplicity (i.e. IRB share model, contracts)
- Participation, engagement and leadership

I.B.4. Patient Experience

- Assess electronic consent process and patient experience
- Evaluate experiences of participating patients

II. Background and Significance

II.A. Significance of Aspirin Dosing: A Global Perspective

ASCVD that leads to ischemic events represents the leading cause of death, morbidity, and disability. Despite remarkable progress in prevention and treatment for atherosclerosis, ASCVD is expected to be an even more prominent cause of death and disability over the next 30 years. In high-income countries, the major factors contributing to this expansion are the aging of the population coupled with increases in incidence of obesity, diabetes, and sedentary lifestyle. Despite declining age-specific disease rates, the total disease burden increases as ASCVD eventually affects a larger population of older adults. In economically developing countries, a major epidemic of atherosclerosis is occurring, concentrated in younger age groups and presumably due to increasing tobacco use as well as obesity and diabetes arising from Westernization of diets and lack of exercise.
The development of new biological and technological approaches to treating ASCVD is exciting, but maximizing the use of an inexpensive yet effective therapy shows more promise for reducing death and disability on a global scale. Numerous clinical trials have shown the clinical benefit of aspirin vs placebo in reducing vascular events in patients with a history of ASCVD or a specific cardiovascular event, but the best dose of aspirin for the general population with ischemic heart disease has not been determined. Considering the burden of ASCVD and that the population affected by it is growing rapidly, identifying the optimal aspirin dose will save lives and prevent ischemic and bleeding events at a global scale.

For example, based on recent evidence suggesting a reduction in ischemic events with lower doses of aspirin, the odds ratio for an event with an aspirin dose of 81 mg/day vs 325 mg/day would be 0.84 (95% confidence interval [CI], 0.64–1.1).13–15 If the rate of death, MI, or stroke in a prospective clinical trial over ~18 months of treatment was 8% with 325 mg of aspirin (based on contemporary trials of aspirin use in patients with ischemic heart disease),13–16–17 then the expected event rate with 81 mg would be 6.8% (95% CI, 5.3–8.7), or ~12 events prevented for every 1000 patients treated. Given the magnitude of the global burden of ischemic heart disease, a 1.2% absolute reduction in events achieved simply through optimal aspirin dosing would be of tremendous importance to public health.

Until recently, aspirin dosing patterns after acute MI in the United States were uncertain. A 2014 analysis of the National Cardiovascular Data Registry’s (NCDR’s) Acute Coronary Treatment and Intervention Outcomes Network (ACTION) Registry-Get with the Guidelines (GWTG) examined aspirin dosing in 221,199 patients with acute MI (both ST-elevation MI and non-ST-elevation MI) from 525 US hospitals.18 Between January 2007 and March 2011, 61% of patients were discharged on 325 mg of aspirin, 36% on 81 mg, and 4% on other doses. The rate of use of 325 mg of aspirin at discharge was 73% in patients who underwent PCI vs. 45% in patients managed medically (i.e., without invasive revascularization). When aspirin was used concomitantly with thienopyridine and warfarin, a 325-mg dose was used in 44% of patients. Even among patients who experienced major in-hospital bleeding, 57% received the 325-mg dose. The relatively high rate of use of this dose, even in patients at high risk of bleeding, and the 25-fold variation in the rate of use of the 325-mg aspirin dose across participating centers are surprising and likely reflect uncertainty regarding appropriate aspirin dosing.18

Further details on aspirin dosing patterns and the impact of high- vs. low-dose aspirin among patients with acute MI undergoing PCI in the United States from 2010-2012 were recently published from the TRANSLATE ACS registry.19 Among 10,213 patients, 6,387 (62.6%) received high-dose (325 mg) aspirin at hospital discharge with substantial variability across the 228 hospitals in the analysis (median hospital-level frequency of high-dose aspirin use was 70%). The adjusted risks of ischemic outcomes (death, MI, stroke, or unplanned revascularization) and bleeding requiring hospitalization through 6 months were similar with high- vs. low-dose (81 mg) aspirin. However, approximately 35% of patients discharged on high-dose aspirin were switched to low-dose aspirin within 6 months. These non-randomized findings, coupled with the findings from the ACTION Registry-GWTG analysis, highlight the substantial variability in aspirin dosing patterns in the United States for patients with ASCVD who have experienced a recent acute MI event and directly point to the need to do an adequately powered, large-scale trial of low- vs.
high-dose aspirin to determine the most effective dose of aspirin for the secondary prevention of ASCVD.

In the United States, the 2010 death rates attributable to coronary heart disease, stroke, and other cardiovascular diseases were 113.6, 39.1, and 82.7 per 100,000, respectively. Globally, given the rapidly increasing burden of ASCVD and limited healthcare resources, particularly in lower-income countries, a similar benefit from identifying the best dose of aspirin for treating the general population with ischemic heart disease could translate to as many as 88,800 fewer deaths from ASCVD annually and would prevent ~145,000 deaths in 2020. In the United States alone, this would mean ~19,000 fewer deaths and MIs each year without employing new treatments or technology and with no additional healthcare expenditures.

In addition to defining the best dose of aspirin from the population perspective, the subgroup analyses and model-based analyses of heterogeneity of treatment effect planned for this proposed trial will allow further insights into refinement of aspirin dosing at the patient level. Such knowledge could further enhance the benefit derived from aspirin treatment.

II.B. Optimal Aspirin Dosing in the Context of PCORnet: A New Model

Although the primary aim of this study is to determine the optimal dose of aspirin for secondary prevention of ASCVD, it also represents the initial use of a transformative approach to developing a new and efficient interactive model for designing and implementing clinical trials that aim to compare the effectiveness of therapies already in use in clinical practice (comparative-effectiveness research, or CER) using methods centered on the needs and experiences of patients. Because we live in an era in which the number of effective (or potentially effective) therapies far exceeds our ability to evaluate them in prospective clinical trials using current methods, we face an urgent need to develop an approach to CER trials that can greatly reduce the cost of trials while maintaining the quality, reproducibility, and generalizability of the research. By using existing data from EHRs organized into the CDM developed by PCORnet and derived from the FDA’s Sentinel project, the trial will develop initial experience with the use of the CDM to supplant costly and time-consuming data collection approaches that are used with traditional clinical trials. By following the majority of patients on the Internet and collecting minimal data directly from them, we can avoid the costs incurred by non-clinically indicated research visits, lengthy case report forms (CRFs), and extensive site management operational approaches.

We believe that a more efficient and less expensive model for trials can be developed that could be extended to more experimental comparisons. However, working through the issues of informed consent, data validation, events ascertainment, and compliance assessment will require acceptance of novel approaches to statistical sampling and “quality by design” principles as well as communication with patients. Amid increasing concerns about patient privacy and research integrity, such an approach to trial efficiency would be difficult to pilot with untested therapies. Because this trial will test only doses of aspirin that are considered relatively safe and are widely used in current clinical practice, it presents a critically and globally important clinical issue with which to develop these new methods.
A trial designed to use existing data resources almost exclusively (supplemented by Internet interaction with research participants and telephone contact from the DCRI Call Center for those without internet access) offers many potential benefits. For example, clinicians will not be burdened with extensive data collection forms and cumbersome consent and contracting procedures. In addition, the patient portal will serve as the primary mechanism for follow-up, with routine data entry by the patients themselves providing a concise set of patient-reported outcomes (PROs).

For physicians and other clinicians, application of the PCORnet CDM format and the patient portal to clinical trials could broaden awareness and participation and, at the same time, aid in conduct of the study. For example, trial enrollment and follow-up could be automated with use of the CDM and patient portal at any time, eliminating or greatly reducing the need for traditional methods such as telephone or postal mail contact, or extended in-person clinic visits. Instead, valuable clinician-patient interaction time can be focused on clinical care and answering questions that arise.

Finally, the platform created by this trial will unite a broad and diverse community of patients and their physicians around the common goal of refining the evidence underlying an existing therapy (aspirin) to maximize its benefit relative to risk. By integrating direct physician and patient participation in the examination of the relationships among clinical outcomes in response to various aspirin doses, this platform will also produce far greater global benefit than the introduction of many other “high-tech” approaches.

II.C. Potential Impact of Proper Dosing of Aspirin

II.C.1. Benefit of Aspirin as Preventive Therapy
Aspirin has a significant impact on the risk of vascular events in patients with known atherosclerosis. The Antiplatelet Trialists’ Collaboration reported a 20%—40% reduction in the risk of death, MI, or stroke for study participants taking aspirin. This benefit was clear for patients with coronary artery disease or cerebrovascular disease and for those with MI, unstable angina, transient ischemic attack (TIA), or stroke. More recently, aspirin therapy was associated with reduced long-term mortality among 6,174 patients undergoing stress echocardiography to evaluate known or suspected ASCVD (hazard ratio [HR] over ~3 years, 0.67; 95% CI, 0.51–0.87; \( P=0.002 \)). After adjustment for the propensity to use aspirin and other possible confounding variables, aspirin use remained associated with a lower risk of death (HR, 0.56; 95% CI, 0.40–0.78; \( P<0.001 \)). Patient characteristics associated with the greatest reductions in mortality included advanced age, known ASCVD, and impaired exercise capacity. These results were recently replicated in a broad assessment of aspirin effectiveness across racial, ethnic, and sex subgroups.

II.C.2. Aspirin Dose and Clinical Outcomes
Despite uncertainties about optimal aspirin dosing, consensus has developed regarding recommendations for aspirin dosing between 75–325 mg daily in patients with ASCVD. This practice has been driven primarily by commercial availability, physician preference, and concerns about adverse effects such as abdominal discomfort and gastrointestinal bleeding associated with higher doses.
Although this dose range has developed empirically over time, attempts to balance clinical benefit with adverse side effects largely reflect indirect comparisons performed in various clinical settings.

Few direct comparisons of different doses of aspirin have been performed, and their results have been inconclusive. The only large, direct prospective study performed to date (OASIS 7-CURRENT)evaluated outcomes among patients with Acute Coronary Syndromes (ACS) over the first 30 days only. The results were complex because the trial used a factorial design of high- and low-dose aspirin with high- and low-dose clopidogrel starting at the time of the index ACS event.Although the factorial analysis showed the most favorable outcomes for the combination of high-dose aspirin and high-dose clopidogrel, the results were not definitive, and a variety of different interpretations have been offered by experts.

Thus, while indirect evidence exists for dose-dependent efficacy of aspirin in preventing vascular events, it is equally clear that no adequately sized randomized trials have addressed this issue, particularly in patients with established ASCVD who are receiving long-term treatment for secondary prevention. The suggestive (but not definitive) data supporting lower aspirin dose emphasize a clear need for larger randomized studies of aspirin dosing in ASCVD.

Various aspirin doses, even as low as 30 mg/day, have been shown to be effective in preventing vascular events. A trial of unstable angina patients found a dose of 75 mg/day to be effective in reducing recurrent vascular events, and the European Stroke Prevention Trial found a benefit of 25 mg twice daily in preventing stroke or death in high-risk patients.

Varied trial results have underscored the uncertainty about aspirin dosing in secondary prevention. A study of secondary prevention after TIA compared 300 mg/day with 1200 mg/day and found no difference in efficacy. Similarly, the Dutch TIA prevention study found no difference in efficacy between 30 and 283 mg of aspirin per day. A trial comparing aspirin doses after carotid endarterectomy found a lower risk of the composite of death, MI, or stroke with daily doses of 81 or 325 mg/day versus 650 or 1300 mg/day. This study contradicted earlier findings of a lower event rate with doses of ≥650 mg versus ≤325 mg per day for prevention of perioperative stroke.

In what had been the largest experience for many years, the Antiplatelet Trialists’ Collaboration’s systematic review of 11 trials of antiplatelet therapy, there was no apparent dose-response relationship of aspirin in secondary prevention of cardiovascular outcomes. When the investigators expanded their meta-analysis to include subsequent trials of antiplatelet therapies, they found that compared with no aspirin therapy, high doses of aspirin (500–1500 mg/day) were not clearly more effective in reducing ischemic vascular events than doses of 75–150 mg/day. Specifically, the proportional reduction in vascular events was 19% with 500–1500 mg/day, 26% with 160–325 mg/day, and 32% with 75–150 mg/day (Table 1).

<table>
<thead>
<tr>
<th>Table 1. Aspirin Dosing and Ischemic Events: ATC</th>
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<tbody>
<tr>
<td>Aspirin Dose</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>500–1500 mg</td>
</tr>
<tr>
<td>160–325 mg</td>
</tr>
<tr>
<td>75–150 mg</td>
</tr>
<tr>
<td>&lt;75 mg</td>
</tr>
<tr>
<td>Any dose</td>
</tr>
</tbody>
</table>
II.C.3. Aspirin and Platelet P2Y<sub>12</sub> Inhibitors
Platelet P2Y<sub>12</sub> inhibitors (ticlopidine, clopidogrel, prasugrel, ticagrelor) reduce the risk of major adverse cardiovascular events (MACE) when added to aspirin (termed dual anti-platelet therapy – DAPT) in patients with ST-segment-elevation myocardial infarction (STEMI), acute coronary syndromes (ACS), and percutaneous revascularization procedures. The recommended duration of DAPT for these indications has been approximately 1 year, but the recently completed DAPT and PEGASUS Trials<sup>25,35</sup> suggest that extended durations of dual antiplatelet therapy for up to 3 years after PCI with coronary stent placement or prior MI provide long-term benefit. Observational comparisons indicate that lower-dose aspirin may be associated with better outcomes when DAPT is used. However, no sizable randomized comparisons of aspirin dosing are available wherein aspirin was used in combination with P2Y<sub>12</sub> inhibitors except for the CURRENT-OASIS-7 trial, as previously mentioned.<sup>25</sup>

Thienopyridine platelet inhibitors have been in clinical use for more than 2 decades. Ticlopidine, the prototype, was shown to reduce ischemic events compared with placebo in a series of clinical trials,<sup>36,37</sup> to be marginally superior to aspirin in one trial in cerebrovascular disease,<sup>38</sup> and to provide additive benefit to aspirin after percutaneous coronary intervention (PCI).<sup>39</sup> Clopidogrel is structurally similar to ticlopidine but is associated with substantially fewer serious adverse events. Most notably, ticlopidine has been associated with the development of neutropenia, thrombocytopenia, and thrombotic thrombocytopenic purpura;<sup>40</sup> such events have not been associated with clopidogrel use in large clinical trials.
The first major study of clopidogrel was the CAPRIE trial, which showed a modest but measurable benefit over aspirin (RR reduction, 8.7%; \( P=0.043 \)) in the secondary prevention of ischemic stroke, MI, or vascular death among patients with vascular disease.\(^1\)

The CURE trial compared the effectiveness of DAPT with aspirin and clopidogrel versus aspirin alone in patients with acute coronary syndromes (ACS).\(^4\) At a mean follow-up of 9 months, there was a 22% relative reduction in the composite endpoint of death, MI, or stroke with combination therapy (9.3% with clopidogrel + aspirin vs 11.4% with aspirin alone; \( P<0.001 \)). Although major bleeding occurred significantly more often with combined therapy (3.7% with aspirin + clopidogrel; 2.7% with aspirin alone; \( P=0.001 \)), there was no excess in major bleeding in the clopidogrel group after the first 30 days, suggesting that most of the risk was related to early revascularization procedures.

Although CURE was not a formal study of secondary prevention (follow-up period was 1 year) and aspirin therapy ranged from 75–325 mg/day, the trial found benefit during 1 year of follow-up. Of note, aspirin dosing in the CURE trial was left to the treating physician’s discretion and was not part of the randomized treatment assignment. With increasing aspirin doses, however, trends toward higher rates of ischemic and bleeding events were observed for patients in both study arms (Table 2), with the lowest rates of bleeding and ischemic events observed in patients taking an aspirin dose of <100 mg/day.\(^4\)

Conflicting results have now been reported for prasugrel and ticagrelor. The TRITON trial compared clopidogrel vs. prasugrel added to aspirin therapy for patients with ACS undergoing PCI and found no evidence for an interaction between aspirin dose and treatment effect of prasugrel relative to clopidogrel for key outcomes, nor did it find a difference in outcomes as a primary function of aspirin dose.\(^4\) In contrast, an observational analysis of the PLATO trial\(^5\) found that lower doses of aspirin were associated with less bleeding and fewer ischemic events in patients receiving ticagrelor. Further, there was a significant interaction between aspirin dose and the treatment benefit of ticagrelor, which overall reduced total mortality compared with clopidogrel. In the PLATO trial, patients receiving low-dose aspirin had a significant reduction in death and MACE if they were randomized to ticagrelor, whereas patients on high-dose aspirin fared equally well with clopidogrel and ticagrelor. However, it should be noted that this comparison of the impact of concomitant aspirin doses with the treatment effect of ticagrelor was non-randomized and was subject to a significant amount of bias based upon regional differences in the concomitant aspirin dose across the multiple countries that participated in the PLATO trial. Nonetheless, the approval of ticagrelor for the treatment of ACS by the Food and Drug Administration (FDA) in the United States incorporated a “black box” warning for avoiding aspirin doses

### Table 2. Effect of Aspirin Dosing on Ischemic Events and Bleeding in the CURE Trial

<table>
<thead>
<tr>
<th>Aspirin Dose</th>
<th>Clopidogrel + ASA</th>
<th>Aspirin</th>
<th>HR (95% CI)</th>
<th>Life-threatening Bleeding</th>
<th>Any Bleeding</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100 mg (n=1927)</td>
<td>80 (8.5%)</td>
<td>96 (9.7%)</td>
<td>0.86 (0.64–1.16)</td>
<td>1.2%</td>
<td>1.9%</td>
</tr>
<tr>
<td>110–161 mg (n=7428)</td>
<td>345 (9.2%)</td>
<td>402 (10.9%)</td>
<td>0.84 (0.73–0.97)</td>
<td>1.7%</td>
<td>2.3%</td>
</tr>
<tr>
<td>&gt;200 mg (n=3201)</td>
<td>157 (9.9%)</td>
<td>221 (13.7%)</td>
<td>0.71 (0.58–0.87)</td>
<td>2.5%</td>
<td>3.9%</td>
</tr>
</tbody>
</table>
>100 mg together with ticagrelor (http://www.fda.gov/NewsEvents/Newsroom/PressAnnouncements/ucm263964.htm).

In summary, the appropriate dose of aspirin in conjunction with a P2Y12 inhibitor is unknown. The ADAPTABLE Trial will investigate this issue by conducting a subgroup analysis according to the concomitant use of a P2Y12 inhibitor and will allow patients to be included in the study who are treated with either clopidogrel or prasugrel, but patients treated with ticagrelor will be excluded given the FDA “black box” warning for the use of high-dose aspirin with ticagrelor.

II.C.4. Systematic Reviews and Mechanistic Insights into Aspirin Dosing

To evaluate the influence of aspirin dosing, a random effects model was used for combining data from 11 randomized, placebo-controlled trials of aspirin in the Antiplatelet Trialists’ Collaboration’s dataset of 14,810 patients. This modeling technique allows adjustments for patient populations and time trends, two limitations inherent in indirect comparisons. Aspirin doses evaluated in this meta-analysis, which examined endpoints including death, acute MI, and stroke (but not bleeding), ranged from 50 mg to 1.5 g daily. Overall, compared with placebo, aspirin resulted in significant reductions in death and death or MI, yet there was a significant decrease in estimated effectiveness as aspirin dose increased (OR for an event: 1.14 with each doubling of aspirin dose; P=0.007). The analysis also suggested that the study population influenced the effect of aspirin, with a greater benefit observed among patients with unstable angina vs MI. For the post-infarction subgroup, the OR was 0.76 (95% CI, 0.66–0.88); for the unstable angina subgroup, the OR was 0.55 (0.43–0.71). Such findings also imply a need for larger randomized comparisons of aspirin dosing in patients with ASCVD.

Although recent observational studies using indirect or nonrandomized comparisons of outcome by aspirin dose suggest that lower doses of aspirin are associated with fewer ischemic events, the possibility remains that in a randomized comparison, higher doses of aspirin may be more favorable. For example, if we consider that aspirin resistance, as defined by elevated urinary thromboxane B2 levels, is associated with worse outcomes and that aspirin 81 mg/day provides less effective thromboxane suppression than aspirin 325 mg/day, we might expect that the latter dose would be associated with better clinical outcomes.

Further, if we postulate that, given its relatively weak antiplatelet effects, an important mechanism of aspirin in prevention of ischemic events is via its anti-inflammatory properties resulting from cyclooxygenase (COX)-2 suppression of platelets or inflammatory cells, similar questions arise about the expected optimal dose. Because 1) aspirin is ~50–100 times more potent in inhibiting platelet COX-1 than monocyte-derived COX-2, and 2) inhibition of COX-2-dependent processes (e.g., inflammation) require larger doses of aspirin (because nucleated cells rapidly resynthesize the enzyme), higher aspirin doses may be required for efficacy. Therefore, if an anti-inflammatory effect of aspirin is important in its ability to prevent ischemic events, then we might expect daily aspirin doses of 162 or 325 mg to be more effective than 81 mg.
In addition, recent research has found alternative pathways for aspirin’s effect on platelet aggregation. These pathways may account for the need for different doses in various individuals and subgroups of patients.

These theoretical and mechanistic arguments highlight the need for a randomized comparison of the effect of aspirin dosing on clinical outcomes that avoids any potential for selection bias toward lower-risk patients receiving lower aspirin doses from their physicians. It also highlights the possible value of mechanistic substudies as a part of this effort to advance our understanding of the factors that identify and modify aspirin response and their relationships with clinical outcomes.

II.D. Aspirin: Mechanism, Clinical Benefit, and Adverse Effects

II.D.1. Mechanism of Aspirin Effect on ASCVD
Aspirin’s putative mechanism of action for preventing cardiovascular events is through irreversible inhibition of platelet COX-1, which prevents the conversion of arachidonic acid to prostaglandin H2—the immediate precursor to thromboxane A2 (TXA2), a potent platelet agonist. Until recently, most experts assumed that the benefit of aspirin solely reflected the inhibition of platelet aggregation. Low-dose aspirin (~80 mg/day) is sufficient to maximally inhibit platelet COX-1 and COX-1-dependent measures of platelet aggregation; higher aspirin doses do not produce additional COX-1 inhibition. Consistent with this finding, selective TXA2 receptor inhibitors have shown no benefit compared with either aspirin or placebo.48,49

With 325 mg/day aspirin dosing, there appear to be additional effects on inhibiting measures of platelet function, particularly from the perspective of non-COX-1 dependent assays (e.g., collagen- and high-dose ADP-induced platelet aggregation). The consequences of this additional, apparently non-COX mediated platelet inhibition with 325 mg/day dosing on clinical outcomes are unknown. However, residual platelet aggregation of non-COX1 dependent platelet aggregation in aspirin treated patients is associated with future cardiovascular events. Further it is clear that additional platelet inhibition in excess of that produced by low-dose aspirin either through inhibition of platelet P2Y12,51 or PAR1,52 receptors can lower the risk of cardiovascular events in high-risk patient populations. In conclusion, aspirin has dose-dependent effects on platelet inhibition that go beyond platelet COX-1 inhibition and which have unknown clinical consequences.

Aspirin is also a weak inhibitor of COX-2, which is expressed in immune function and endothelial cells. Full suppression of COX-2 activity by aspirin has been estimated to require aspirin doses >500 mg/day. Beyond COX1/2 inhibition, it is increasingly clear that aspirin possesses several additional properties that do not appear to depend on COX inhibition. For example, aspirin (and its stable metabolite salicylate) are well-known to produce dose-dependent inhibition of elements of the proinflammatory NFkB pathway: NFkB,54,55 IκB phosphorylation,56 IκB kinase,57 as well as other cellular kinases.58,59 Aspirin also influences erythrocytes,60 endothelial cells,61 endothelial progenitor cells,62 and inflammatory markers (CRP, interleukin-6, and macrophage colony stimulating factor (MCSF).63 In addition, omics-based assays are now illuminating the full suite of pathways affected by aspirin, many of which are not predicted
based on aspirin’s putative mechanisms of action: acetylation of proteins beyond COX, alterations in fatty acid and amino acid metabolism, and changes in platelet RNA/protein content. Although it is unknown to what extent these apparent non-COX effects of aspirin are dose-dependent, it is clear that aspirin interacts with a wide array of diverse biologic pathways that may underlie its beneficial effects on cardiovascular disease.

In contrast, aspirin may also interact with prothrombotic pathways. In addition to serving as a precursor to TXA2, PGH2 produced by COX1/2 can be metabolized by vascular endothelial cells to produce prostacyclin (PGI2). Although TXA2 promotes platelet aggregation and vasoconstriction, PGI2 inhibits platelet aggregation and induces vasodilation. Aspirin causes dose-dependent reductions in TXA2 and PGI2 production, thus producing a blend of anti- and prothrombotic effects. No studies to date have directly established that more profound suppression of PGI2 formation by higher doses of aspirin is sufficient to initiate or to predispose to thrombosis. However, recent studies in healthy volunteers exposed to aspirin 325 mg/day demonstrate that in certain individuals such dosing can lead to a paradoxical increase in platelet aggregation despite complete suppression of platelet COX-1. Further, recent studies have associated selective COX-2 inhibitors with unopposed platelet activation and a greater likelihood for thrombotic events. These findings provide mechanistic support for the efficacy of lower aspirin doses in preventing COX-1-mediated TXA2 production while preserving the antiaggregatory and vasodilatory effects of PGI2. Whether higher aspirin doses will achieve a more beneficial (or potentially harmful) balance of TXA2 vs PGI2 in preventing cardiovascular events is unknown.

II.D.2. Aspirin Intolerance and Dose-Related Risks of Aspirin Therapy

In patients with established ASCVD, the risk associated with aspirin use is quite low compared with the benefit. Only a very small number of patients develop a serious anaphylactic reaction or bronchoconstriction with nasal polyps when they take aspirin. The risk of intracranial hemorrhage with chronic aspirin treatment is estimated to be <0.04% per year. A modest number of patients develop serious gastrointestinal bleeding with chronic aspirin therapy, mostly related to the loss of the protective effect of prostaglandins on the gastric mucosa. A larger number of patients have gastrointestinal intolerance with aspirin, but this is often transient and can be overcome in many patients. The absolute risk varies as a function of the trial entry criteria, but none of these major risks exceeds 5 events per 100 patients treated per year. Except for patients with pre-existing asthma, no studies have described the risk factors for these complications of aspirin use.

The Clopidogrel versus Aspirin in Patients at Risk of Ischaemic Events (CAPRIE), Coumadin Aspirin Reinfarction Study (CARS), and Sibrafiban versus aspirin to Yield Maximum Protection from ischemic Heart events post acute coronary syndromes (SYMPHONY) trials have provided excellent opportunities to understand the incidence of aspirin intolerance and toxicity in patients with vascular disease. Among patients randomized to aspirin therapy alone in CARS, 2.7% discontinued the drug permanently and 7.6% had a dose reduction. In SYMPHONY, which compared aspirin 80 mg twice daily with high and low doses of the oral glycoprotein IIb/IIIa inhibitor sibrafiban, the rate of early aspirin discontinuation was 19.2%, although over half of these patients later reported open-label use. In CAPRIE, 11% of patients randomized to aspirin therapy had to stop the drug because of rash, diarrhea,
indigestion, minor or major bleeding, or abnormal liver-function tests (Table 3). Much higher proportions of patients had minor adverse events that did not lead to discontinuation.

| Table 3. Adverse Events with Antiplatelet Therapy in the CAPRIE Trial |
|--------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                         | Clopidogrel | Aspirin | Clopidogrel | Aspirin | Clopidogrel | Aspirin |
| Rash                    | 578 (6.0)   | 442 (4.6)* | 25 (0.26)   | 10 (0.1)* | 86 (0.90)  | 39 (0.41)* |
| Diarrhea                | 428 (4.5)   | 322 (3.4)* | 22 (0.23)   | 11 (0.1)  | 40 (0.42)  | 26 (0.27)   |
| Indigestion/N/V         | 1441 (15)   | 1686 (18)* | 93 (0.97)   | 118 (1.2) | 182 (1.9)  | 231 (2.4)*  |
| Any bleeding disorder  | 890 (9.3)   | 890 (9.3)  | 132 (1.38)  | 149 (1.6) | 115 (1.20) | 131 (1.4)   |
| Intracranial hemorrhage | 34 (0.35)   | 47 (0.49)  | 30 (0.31)   | 41 (0.43) | 20 (0.21)  | 32 (0.33)   |
| GI hemorrhage           | 191 (2.0)   | 255 (2.7)* | 47 (0.49)   | 68 (0.71)*| 50 (0.52)  | 89 (0.93)*  |
| Abnormal liver function | 285 (3.0)   | 302 (3.2)* | 11 (0.11)   | 9 (0.09)  | 22 (0.23)  | 28 (0.29)   |

* P < 0.05. Data are number (%) of patients. N/V = nausea/vomiting; GI = gastrointestinal.

Although the dose-dependent COX inhibition by aspirin suggests that gastrointestinal bleeding can be reduced with lower aspirin doses, no large, prospective randomized study has examined the relationship between lower doses of aspirin and bleeding. The available evidence is equivocal, and much of the available evidence is from an era in which positive reporting bias was common, especially in observational studies.

Despite the limitations of the available literature, a dose-response relationship has been suggested with regard to GI side effects, including discomfort and bleeding. The lower risk associated with lower doses is believed to reflect both the differential inhibition of COX-1 on platelets and the differential inhibition of COX-2 in the gastric mucosa. In a case-control study, the odds ratio for hospitalization for a bleeding peptic ulcer was 2.3 with 75 mg of aspirin, 3.2 with 150 mg of aspirin, and 3.9 with 300 mg of aspirin. In another study, higher doses (300–1200 mg/day) showed a consistent dose-response relationship between gastrointestinal bleeding and a need for hospitalization. Conversely, a systematic review that included data from 66,000 patients reported no difference in the incidence of gastrointestinal hemorrhage with doses <163 mg/day vs higher doses. This study did not examine the risk of bleeding with doses <100 mg/day relative to higher doses, which may be important, because prostaglandin synthesis is inhibited with doses >100 mg.

There is no reliable estimate of the risk of fatal gastrointestinal bleeding as a function of aspirin doses below 162 mg/day. A retrospective analysis of antiplatelet therapy in the CURE trial implies lower risks of both life-threatening bleeding and overall bleeding with lower aspirin doses (see Section II.C.3). The risk of death, in the presence of clinically detectable bleeding from a gastric ulcer, is estimated to be 0.5%–10%. Considering the available evidence, aspirin doses <100 mg/day may well be associated with less gastrointestinal bleeding, which may in turn translate into a decrease in fatal events.
A large systematic review in 2007 found no evidence that doses higher than 81–100 mg/day were better for ischemic events and lower doses seemed to be associated with less bleeding. However, despite the millions of people taking aspirin daily, fewer than 5,000 participants in randomized trials could be aggregated, leaving the analysis dependent on extrapolations and observational studies.

In summary, more than 16 million Americans have ASCVD, contributing to substantial morbidity, mortality, and costs. Based on the side-effect profile reported in trials of aspirin therapy, 3% of these patients would be expected to have a total of 319,000 episodes of gastrointestinal bleeding at an aspirin dose of 325 mg/day. Even assuming a conservative 10% reduction in gastrointestinal bleeding with optimal dosing of aspirin, about 32,000 such bleeding events and 3,200 deaths from gastrointestinal hemorrhage would be prevented annually in the United States alone. Accordingly, there is a clear need for an adequately powered randomized controlled trial to define the optimal dose of aspirin in order to minimize bleeding risks while preventing ischemic events.

II.E. Modifiers of Aspirin Dose

II.E.1. Aspirin, Minorities, and Other Subgroups
Early trials with aspirin were performed almost exclusively in men, leading to the empirical observation that the significant reductions in events were limited to men. Further study showed this simply was a problem of inadequate power to detect an effect in women, because very few women had been enrolled in trials. In a large prospective cohort study of 28,678 nurses taking one to six aspirins (dosage unknown) per week, the age-adjusted relative risk (RR) of a first MI was 0.68 (P=0.005) with a trend toward fewer cardiovascular deaths (RR, 0.89; P=0.56). Still, without randomized trials enrolling representative numbers of men and women, it remains unknown whether aspirin has the same effect in men and women or if it shows a similar relation between dose and outcome.

In addition, little is known about how the effect of aspirin is modulated by age, race and ethnic background, the presence of diabetes or renal function. Because ASCVD is the leading cause of death in women, racial minorities, diabetic patients, and patients with CKD in the United States, and because its prevalence and associated adverse events increase proportionately with age, it is critically important to perform randomized trials of aspirin use in these populations. Patients with diabetes constitute a particularly important subgroup, given the considerable evidence that they may exhibit resistance to the antiplatelet effects of aspirin.

II.E.2. Other Issues

II.E.2.1. Enteric Coating
Enteric coating of aspirin became popular as a proposed approach to reducing aspirin’s gastrointestinal toxicity. While some small studies have shown promising results regarding the benefit of enteric coating, no large outcomes trials have evaluated the benefit-risk balance of enteric-coated aspirin versus aspirin without enteric coating. Additionally, pharmacokinetic/pharmacodynamic studies have raised questions about the reliability of absorption of aspirin when given in enteric-coated form. Given these uncertainties, we will collect details of the actual dose and type (enteric-coated vs. non-
enteric-coated) of aspirin taken by trial participants at regular intervals, but the randomization to low vs. high-dose aspirin will not specify whether the patient should take enteric-coated aspirin.

III. Research Design and Methods

III.A. Study Aims 1 to 3: Aspirin Dosing
We plan to compare the effectiveness of two once-daily doses of aspirin (81 mg and 325 mg) in a secondary-prevention trial in patients with ASCVD, using a novel format that exploits EHR data that have been standardized according to a common format and primarily web-based systems of communication among enrolled patients and trial investigators with the support of health systems interested in the best care for their patients. Every aspect of the trial is designed to not only answer the research question, but also to build an infrastructure for future pragmatic trials in which a community of patients, clinicians, researchers and administrators work together to improve patient care and clinical outcomes.

III.A.1. Enrollment and Eligibility
Importantly, these criteria are intended to reflect the best judgment of clinicians in practice and to reflect the general “uncertainty principle.” For patients whom aspirin is indicated to reduce recurrent events, the clinician is uncertain about the best dose of aspirin to prescribe. These enrollment criteria have been selected to achieve the most generalizable sample possible to address the main study hypothesis. Furthermore, the protocol was posted for public review and comment during July, 2015. Specific questions regarding key eligibility criteria were voted upon and the responses are described herein. Approximately 57% of the survey respondents voted to not limit the inclusion criteria to only include patients already taking aspirin at the time of randomization (with 7% expressing no opinion), 49% voted to allow patients taking ticagrelor at the time of screening to be included in the trial (with 14% expressing no opinion), and 49% voted to exclude patients with a clear indication for an oral anticoagulant even if they are not currently taking an oral anti-coagulant (with 6% expressing no opinion). The final inclusion/exclusion criteria listed below were adapted based upon these survey responses, but based upon feedback from the CDRNs regarding their local Institutional Review Boards’ perspective on the “minimal risk” of the informed consent process, it was decided to exclude patients treated with ticagrelor given the “black box” warning in the ticagrelor label in the United States for avoiding high-dose aspirin with concomitant ticagrelor use.

1. Known atherosclerotic cardiovascular disease (ASCVD), defined by a history of prior myocardial infarction, prior coronary angiography showing ≥75% stenosis of at least one epicardial coronary vessel, or prior coronary revascularization procedures (either PCI or CABG)
2. Age ≥ 18 years
3. No known safety concerns or side effects considered to be related to aspirin, including
   a. No history of significant allergy to aspirin such as anaphylaxis, urticaria, or significant gastrointestinal intolerances
   b. No history of significant GI bleed within the past 12 months
c. Significant bleeding disorders that preclude the use of aspirin

4. Access to the Internet. In the event that the CDRNs are notified that a cohort of patients without internet access can be included, then patient agreement will be obtained during the consent process to provide follow-up information by telephone contact with the DCRI Call Center.

5. Not currently treated with an oral anticoagulant – either warfarin or a novel anticoagulant (dabigatran, rivaroxaban, apixaban, edoxaban) – and not planned to be treated in the future with an oral anticoagulant for existing indications such as atrial fibrillation, deep venous thrombosis, or pulmonary embolism.

6. Not currently treated with ticagrelor and not planned to be treated in the future with ticagrelor.

7. Female patients who are not pregnant or nursing an infant

8. Estimated risk of a major cardiovascular event (MACE) > 8% over next 3 years as defined by the presence of at least one or more of the following enrichment factors:
   a. Age > 65 years
   b. Serum creatinine > 1.5 mg/dL
   c. Diabetes mellitus (Type 1 or Type 2)
   d. 3-vessel coronary artery disease
   e. Cerebrovascular disease and/or peripheral arterial disease
   f. Left ventricular ejection fraction (LVEF) < 50%
   g. Current cigarette smoker

There will be no exclusions for any upper age limit, comorbid conditions, or concomitant medications other than oral anticoagulants and ticagrelor that are used at the time of randomization, or are planned to be used during the study follow-up.

Simple, inclusive eligibility criteria will make enrollment easier, and will render study results more generalizable to a broader population of patients. We will exclude pregnant or lactating women (because of concern for the fetus or child), patients taking oral anticoagulants or likely to require an oral anticoagulant during trial follow-up (because of complex drug interactions and a projected excessive risk of bleeding), and patients at relatively low risk for cardiovascular events (ie, no enrichment factor because of the large number of outcomes needed to detect a clinically meaningful difference with the available sample size).

III.A.2. Solicitation of Participation
The majority of the PCORnet’s Clinical Data Research Networks (CDRNs) and 1 Patient-Powered Research Network (PPRN) have agreed to participate in this trial. After obtaining IRB approval of the trial and with agreement of the clinicians and their health systems, the EHRs at the participating health systems in the CDRNs will be queried for eligible study participants. This will be supplemented with local engagement of health systems, clinicians, and patients as well as direct recruitment in clinics and hospitals. The trial is designed not to interfere with routine clinical practice and is expected to impose a minimal burden on clinicians, clinics, health systems, and patients.
III.A.2.a. Women, Elderly, and Minority Subjects

The inclusion of women and minorities in clinical trials is desirable for scientific, ethical, and social reasons. This trial’s investigators feel strongly that the inclusion of women, the elderly, and racial minorities is imperative to meet the goal of developing a therapy that is ultimately generalizable to the global ASCVD population. The increasing number of men and women ≥65 years of age with ischemic heart disease also mandates that they be represented in a dosing trial evaluating the safety and effectiveness of aspirin. Although octogenarians represent only 5% of the U.S. population, they represent 20% of all patients hospitalized for MI and 30% of all MI-related hospital deaths.84

PCORnet is committed to promoting equity in research and PCORI has made concerted efforts to include underrepresented populations in research. Because this trial is intended to identify an optimal aspirin dose that is applicable to a large population of ASCVD patients, our aim is to enroll a study population whose demographic profile is similar to the representation of women, elderly persons, and minorities in the overall population of individuals with ASCVD in the United States.20,85 Estimates of the incidence of ASCVD in American women have ranged from 30%86 to 51%85 and vary considerably relative to age group.87 Although the elderly (>75 years) account for <6% of the population,88 the incidence of ASCVD among elderly patients is proportionally higher compared with younger age groups (169 vs. 60 per 1,000 persons).88 Based on these demographics, we will aim for 35%-40% of our enrolled patients to be women and at least 10% to be elderly (≥75 years). Because the incidence of ASCVD is similar among ethnic groups in the United States (~7% for each group of non-Hispanic whites, non-Hispanic blacks, and Hispanics),85 we have selected to strive for enrollment target of 20-25% for minorities to reflect their representation in the general population (e.g., 12.1% non-Hispanic blacks, 12.3% Hispanics).88

The CDRNs will develop local recruitment strategies specifically designed to facilitate the inclusion of a broadly representative population of patients that incorporates women, the elderly, and racial/ethnic minorities. For example, many participating sites are regional centers located in areas with higher numbers of minorities; this will enhance minority recruitment.

III.A.2.b. Sources of Potential Bias

Because this trial emphasizes the use of the Internet and standardized EHR data, we acknowledge potential barriers to participation including older age, low socioeconomic status, and low literacy. However, Internet use will facilitate the conduct of this pragmatic trial by reducing potential limitations to enrollment, including the costs associated with frequent follow-up visits at healthcare facilities. We will encourage enrollment of patients with personal Internet access as well as patients with public Internet access only, such as those available through a library, enrolling physician’s office, or workplace. Additionally, the ADAPTABLE patient portal will be accessible via tablet computers and smart phones so patients with access to the internet only through cellular service will be able to participate in this manner.

No studies have formally evaluated the association of Internet use with general health status and outcomes. Internet users may derive indirect benefits that improve their health status (e.g., disease recognition, symptom awareness, risk-factor modification), and/or they may be better educated, more
motivated in personal health maintenance, and wealthier, all of which may be associated with better health status and outcomes in secondary prevention. Although the investigators acknowledge these possibilities, we believe the results of this trial remain applicable to all patients with ASCVD for three reasons.

First, since the occurrence of cardiovascular events is multifactorial, standard risk factors (e.g., smoking, diabetes, and age) are more likely to directly influence cardiovascular event rates than Internet use. Second, aspirin use has substantial consequences, effecting a ~25% reduction in adverse cardiac events. Rather than modify the benefit of aspirin therapy, Internet use would more likely be associated with increased awareness of the general need for aspirin therapy, which also is an educational objective of the trial patient portal. Third, the aim of this trial is to define the optimal dose of aspirin in patients with ASCVD. Even if Internet use were associated with improved health, this effect would be a general reduction in adverse events across all treatment groups. Although attributing a global improvement in outcomes in this trial to Internet use may be hypothesis-generating, such a finding still would not influence the primary objective of the trial.

Another source of bias may exist with respect to aspirin therapy prior to randomizations. We expect that most patients recruited into the ADAPTABLE trial will already be treated with long-term aspirin therapy, given the nature of the inclusion criteria and the known high utilization of aspirin for the secondary prevention of coronary artery disease in the United States. However, we will add a sensitivity analysis with a 10-day landmark to the statistical analysis plan (SAP) that excludes all events occurring in the first 10 days following randomization in order to formally test for a legacy effect of the aspirin dose taken before randomization.

**III.A.2.d. Patient Engagement**

The ADAPTABLE trial will address several key questions mentioned in PCORI's definition of patient-centered outcomes research. This trial is designed to help individuals answer the following questions: "Given my personal characteristics, conditions, and preferences, what should I expect will happen to me?" "What are my options and what are the potential benefits and harms of those options?" "What can I do to improve the outcomes that are most important to me?"

The study seeks to make more relevant, detailed information regarding aspirin therapy as secondary prevention for ASCVD available to patients and their caregivers by answering question such as “What is the most effective dose of aspirin in secondary prevention of ASCVD?” and “What are the bleeding risks associated with taking 81mg or 325mg of aspirin?” Of the forty topics suggested for the first PCORnet study, six were selected for consideration through the PCORI Advisory Panel Prioritization process by a body consisting of patients, scientists, and other stakeholders charged with prioritizing CER topics for PCORI. ADAPTABLE was chosen to be the first PCORnet study based on its importance to patients, clinicians and scientific communities.

Patient and stakeholder engagement has been a priority to the study team since the inception of ADAPTABLE. A partnership was formed with the Health eHeart Alliance Patient-powered Research
Network to support ADAPTABLE by leading the CDRN patient representative team called the Adaptors. Additionally, two seasoned cardiology patient advocates served on the protocol design committee to assist with defining essential characteristics of the study, participants, and outcomes. These advocates also worked actively with the study team and Health eHeart to develop the patient engagement plan.

The DCRI Coordinating Center for ADAPTABLE will convene the Executive Committee, Steering Committee, and Data and Safety Monitoring Board (DSMB) – each of which will have at least 2 patient representatives, while Health eHeart will support and facilitate the Patient Group herein referred to as “Adaptors.” The Adaptors group will include one patient representative from each participating CDRN. CDRNs will seek to identify patients who represent the trial’s population (see Appendix: Patient Survey), and they may select skilled cardiology patient representatives who use aspirin if they are unable to locate a patient representative with appropriately matched characteristics and skills. Diversity and inclusiveness are highly valued and encouraged. We will ensure that the patient representatives will be recruited from the population they are serving by using the Adaptors Screening Tool (see attachment provided with this document). The screening tool will be circulated by Health eHeart and partnering CDRNs to identify potential patient representatives for the Adaptors group. The Adaptors will serve a dual role as designers and advisors; they will work in concert with the study team to help design study materials, the trial consent form, and recruitment plans through their work with Health eHeart. They will also be members of the Steering Committee where they will monitor study conduct and progress. The Co-Chairs of the Adaptors group, who will be selected by the patient representatives and confirmed by Health eHeart and the DCRI Coordinating Center to be able to carry out the associated duties, will also serve as the patient representatives on the Executive Committee to provide study oversight, promote cross-pollination of ideas and sharing of information between the Executive Committee, Steering Committee and the Adaptors Group. Also, two cardiology patient advocates (not a part of the Adaptors) will serve on the DMSB to participate in review of study data sets and adverse event monitoring and reporting.

The ADAPTABLE Co-Learning Community (ACLC) will be a discussion forum supported by Health eHeart and composed of Adaptor patient representatives, investigators, clinicians, study team members, and support staff from participating networks, and associated professional society and disease advocacy organization representatives. The forum will promote information sharing from diverse perspectives with anticipated discussion of issues related to physician-patient communications, adverse events potentially related to aspirin, common patient questions, adherence, and patient participation in
research. The forum will be co-managed by team members from Health eHeart and the DCRI Coordinating Center to ensure that study-related issues are properly addressed through appropriate channels such as committee meetings. However, a discussion forum is available to promote co-learning, sharing of information and community building in a less formal manner than structured, time-constrained committee meetings. The diagram at right shows how patients will be engaged in all operations of the trial.

III.A.3. Study Design and Procedures

III.A.3.a. General Enrollment Plan and Timeline

To meet the goal of 20,000 patients, we plan to enlist the CDRNs and patient advocates to mobilize the requisite number of clinics and health system based populations. Enrollment of at least 2,500 patients per CDRN over an approximate 24-month period is expected. Informed consent will be required from all participants. The anticipated study duration is 3 years; 3 months for planning, 24 months for enrollment, a maximum of 30 months for follow-up, and 3 months for data analysis.

III.A.3.b. Cohort Identification

Local site investigators within the CDRNs will be asked to endorse the protocol. They will then be asked to give their permission for the CDRN, through its integrated health system members, to identify and contact potentially eligible patients. In the latter case, patients who meet criteria for secondary prevention after a cardiovascular event will be identified using search algorithms developed by the DCRI Coordinating Center (based on the trial inclusion criteria) and customized by the CDRN for their own EHR systems. Broad agreement from both cardiovascular specialists and primary care physicians will be sought. In this trial, we believe that most systems will agree that prior approval of the relevant clinician will be needed and useful since these patients will be at high-risk for death or a major disabling event. Although it is unlikely that a medical reason for ineligibility will be found, most of these patients are close to their clinicians, whose confidence in and support of the trial will be important for patient engagement, both in terms of participation as well as promoting adherence to the study medication and treatment of the inevitable clinical events.

III.A.3.c. Enrollment, Randomization, and Drug Allocation

Patients who are identified as candidates for the trial will be directed to the ADAPTABLE patient portal for the eConsent as well as an abbreviated eligibility confirmation prior to randomization. The ADAPTABLE patient portal will include the electronic patient consent (eConsent) that satisfies the local Institutional Review Board (IRB) and state law requirements of the participating CDRNs. CDRN personnel will be available, if needed, to give a more local perspective on protocol-related consultations.

Patients will be randomized via the patient portal in a 1:1 ratio to receive 81 mg vs. 325 mg of aspirin. Patients will be asked to obtain their randomized aspirin dose at their local pharmacies. The randomization scheme will be established before the inception of enrollment. In a trial of this size, stratified randomization has no significant advantages.
After randomization, patients will be asked to answer questions related to current aspirin use and dose, and other specific concomitant medications. They will also be asked to provide contact information such as their name, email and home address, phone numbers and contact information for a family member or friend not living with them. This data will be used to contact patients that miss multiple visits and to identify events such as hospitalizations that occur out of network.

Neither patients nor health care providers will be blinded to their treatment assignment as blinding would add substantial complexity and cost without commensurate incremental benefit. The “hard” outcomes, large sample size and equipoise in the clinical community should enable valid results to be obtained, and we have no reason to believe that investigator or clinician bias will play a role in ascertainment or classification of events.

**III.A.3.c. Concomitant Therapy**

Key concomitant medications will be self-reported by the participant, and over time will also be harvested from the PCORnet DataMart in CDM version 3.0 as it is deployed. Pre-randomization aspirin dose for aspirin users will be recorded. Participants will report key concomitant medications annually, either on the patient portal or during phone follow-up calls from the DCRI Call Center.

Details on the use of aspirin and important concomitant medications (P2Y12 inhibitors) and other over-the-counter medications of interest (non-steroidal anti-inflammatory agents and proton-pump inhibitors) will be collected from patients at baseline and annually during study follow-up. Details on other common secondary prevention medications prescribed (beta-blockers, ACE inhibitors/angiotensin receptor blockers, and statins) will be collected from the CDM. The large trial population and randomization will minimize potential biases and confounding that could result from differential use of concomitant medications by randomized treatment assignment. A sensitivity analysis that accounts for concomitant medications as time-dependent covariates will be done to account for post-randomization treatments.
### III.A.3.d. Schedule of Assessments

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<th>Baseline All participants</th>
<th>1-3 Week Follow-up Post-Randomization</th>
<th>Every 3 or 6 Month Follow-up</th>
<th>Annual Follow-up (12 &amp; 24 month)</th>
<th>End of Study</th>
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<td>Video describing trial expectations</td>
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<td>Collect demographic/ contact information</td>
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<td>Collect key concomitant (Rx and OTC) medications</td>
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</table>
Ill.A.3.e. Data Collection and Follow-up

For all participants and during follow-up, demographics and key medical history, cardiovascular risk factors, and certain current medications will be obtained from the CDM. Patients will be randomly assigned to follow-up contact every 3 vs. 6 months via the ADAPTABLE patient portal. During follow-up contacts, data on aspirin dose and use, and hospitalizations will be collected. By embedding a randomization for follow-up every 3 months vs. every 6 months for patients via the ADAPTABLE patient portal, we will be able to investigate the best method for optimizing patient follow-up and data collection in this pragmatic trial. In the event that the CDRNs are notified that a cohort of patients without internet access can be included, then these patients will be contacted every 6 months by the DCRI call center via telephone for the follow-up contacts.

As envisioned by PCORnet, the PCORnet distributed research network (DRN) is to be a “…functional distributed research network that facilitates multi-site patient–centered research across the CDRNs, the PPRN, and other interested contributors. The distributed network enables the conduct of observational research and clinical trials while allowing each participating organization to maintain physical and operational control over its data.”

In PCORnet’s distributed data environment, code is developed centrally and distributed to each partner to execute against data that are stored in a common format. Code (“queries”) are distributed and results are returned via PopMedNet, a networking software application that manages the creation, operation, and governance of distributed health data networks.

The PCORnet CDM is the foundation of the PCORnet DRN. The PCORnet CDM is being implemented in phases to allow for the incorporation of new data domains and fields based on PCORnet needs, lessons learned from use, and data availability.

The CDM contains some of the 18 elements that define PHI under HIPAA, including encounter dates and date of birth. The necessary “cross-walks” between the arbitrary identifiers included in the CDM and their originating data are not specified in the scope of the CDM, but are expected to be maintained by each CDRN.

- PATID is a pseudoidentifier with a consistent crosswalk to the true identifier retained by the source site. For analytical data sets requiring patient-level data, only the pseudoidentifier is used to link across all information belonging to a patient.
- Locally maintained “mapping tables” are tables necessary to implement so that each CDRN/partner has the ability to map arbitrary identifiers back to the originating data and patient.

The PCORnet CDM Version 3.0 includes ten tables that represent specific data domains that are available in EHRs and directly from patients. Table 4 describes data categories that are applicable to the ADAPTABLE case report form (CRF):
Table 4. Data Categories Applicable to the ADAPTABLE CRF

<table>
<thead>
<tr>
<th>Description</th>
<th>Applicability for ADAPTABLE Data Collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographic</td>
<td>Birth date, sex, Hispanic ethnicity (y/n) &amp; race captured.</td>
</tr>
<tr>
<td>Encounter</td>
<td>Encounter type will be used to identify hospitalizations during follow-up period. Deaths occurring during a hospital stay will be captured in the discharge status variable.</td>
</tr>
<tr>
<td>Diagnosis</td>
<td>Diagnosis codes and associated encounter dates will be used to establish medical history prior to randomization.</td>
</tr>
<tr>
<td>Procedure</td>
<td>Procedure codes and associated encounter dates will be used to establish cath, PCI, and CABG prior to randomization.</td>
</tr>
<tr>
<td>Vital</td>
<td>Height and weight are captured; tobacco status has been added to v2.0.</td>
</tr>
<tr>
<td>Lab Result</td>
<td>Creatinine, hemoglobin, and LDL-C will be included in v2.0.</td>
</tr>
</tbody>
</table>

In essence, except for aspirin dose and use of over the counter NSAIDS, H2 Blockers and PPIs, all of the essential data for the trial are included in CDM 3.0. A unique attribute of this trial is the continuous improvement expected in the extent and complexity of the available data as the trial goes on.

III.A.3.d.i. Minimizing Cross-Overs

Because this is an open-label trial in which participants and their healthcare providers will know their randomized dose of aspirin, the CDRN will be asked to engage regularly with the healthcare providers in their network regarding the rationale for the trial and the importance of compliance with the randomized treatment assignment to minimize patient cross-over. Health care providers will be asked not to change a patient’s aspirin dose during the trial unless it is absolutely necessary to ensure patient safety. On the ADAPTABLE patient portal, patients will be reminded of the importance of adhering to the aspirin dose they were randomly assigned to unless their physician changes their dose for a major safety concern. Patients who cross-over will be analyzed as they were originally randomized according to the intention-to-treat principle.

III.A.3.d.ii. Drug Discontinuation – Monitoring and Recommendations

Aspirin use will be collected via the responses to the patient questions during follow-up. During each follow-up contact, compliance with the randomized aspirin dose will be assessed. The occurrence of and reasons for discontinuing aspirin or changing the dose of aspirin will be ascertained. In the event of aspirin intolerance or bleeding, the decision to continue the randomized aspirin dose, convert to a lower or higher aspirin dose, or to discontinue aspirin therapy will be left to the treating physician’s discretion. Patients who discontinue aspirin therapy altogether will remain in the trial for clinical outcomes follow-up according to the intention-to-treat principle.

If a female participant becomes pregnant or plans to attempt to become pregnant during study follow-up, then aspirin should be permanently discontinued. Furthermore, if a participant requires an oral anticoagulant during study follow-up, the treating physician should carefully consider whether to continue...
aspirin based upon the estimated bleeding risks of aspirin used together with an oral anti-coagulant based on the clinical context. Finally, participants who develop an intracranial hemorrhage or a life-threatening bleeding event during study follow-up should be considered for permanent discontinuation of aspirin, at the discretion of their treating physician.

**III.A.3.d.iii. Delinquent or Missing Follow-up**

Data from patients who fail to respond to questionnaires via the ADAPTABLE patient portal or via telephone contact with the DCRI Call Center after at least two separate contact time points will be collected using all available approaches including but not limited to the DCRI call center, patient finder companies, the Internet, EHR data stored in the CDM format in the PCORnet DataMart, the Social Security Death Index, and other additional search methods. Additionally, if these methods fail, the site will be contacted to determine whether they have been in contact with the patient.

**III.A.3.e. Developing and Refining the PCORnet Infrastructure**

Based on previous experience with clinical trials and patient follow-up (see Section III.A.3.d), there is broad familiarity with the potential limitations of EHR data and patient-reported clinical outcomes. As a result, PCORNet is in a position to identify and resolve inaccuracies to improve the validity of such data. Because this trial will investigate a clinically relevant issue in a new format, the study will also examine the potential for conducting trials through health systems with secondary data use and Internet-based follow-up with both participants and clinician-investigators. Because this trial will rely in large part on patient-reported data gathered via the Internet, it will enable us to show the feasibility of participant enrollment and reporting via the Internet. As previously stated, Aim 3 of this project will focus upon using defined metrics to evaluate the PCORnet infrastructure that will be developed across PCORnet studies.

**III.A.3.f. ADAPTABLE Patient Portal**

Patient data will be captured via the ADAPTABLE patient portal that will be designed and maintained by Mytrus. Screens will be user-friendly and will contain questions that are easy to understand and answer. Content for the Patient Portal will be in English and Spanish to facilitate bi-lingual interactions. Based on previous studies of Internet usability for people >65 years, the study patient portal will be designed to accommodate issues such as impaired vision (e.g., font size and “readability”), memory (e.g., focused error messages and hypertext links), and motor control and precision of movement (e.g., “clickability”). Screens, checkboxes or pull-down menus will be used when possible for data entry, to minimize the need to enter free text.

Queries will be programmed directly into the study patient portal and will prompt patients for missing or discrepant data. For example, if a patient skips a question, a program warning will occur that directs the patient to complete all questions before continuing. Similarly, program warnings will alert patients if out of range data have been detected. This method of on-the-spot querying will ensure cleaner data.

The patient portal will contain a general information section that allows access by study participants and general and medical lay communities. There will also be restricted sections on the patient portal that are
accessible only to the trial’s investigators. The restricted section for investigators will contain more specific information about their local participation. Visibility of the link to the restricted section will be determined by the access rights.

Below is a partial list of items that could be shared:

- **Training materials**, to include resources for training in ethics of human research, recordings of investigator meetings/conference calls, and other appropriate training materials.

- **Educational links**, to provide additional education materials for patients (e.g., the American Heart Association website) and providers (e.g., theheart.org; the American College of Cardiology website).

- A **protocol** section, to include the current version of the protocol. Eligibility criteria will be listed, as will instructions for completing follow-up questionnaires through the patient portal.

- Regular communications, to include enrollment data, frequently asked questions (FAQs), and other trial information.

- **Enrollment information** to be posted on the general area of the site.

### III.A.4. Data Security and Back-up Procedures


The system is deployed using a LAMP (Linux, Apache, MySQL, PHP) technology stack running on a virtual machine (VM) in a private cloud. By using a virtualized environment, additional resources (disk space, processing power) can be added to the VM as necessary to support increased load. Data centers used to host the system hold current SSAE16 SOC certification and have Tier 1 internet access from multiple major Internet Service Providers (ISPs) for redundancy and 100% network uptime.

Access to the system’s web-based user interface (UI) is provided using HTTPS (web protocol that is secured using 128-bit [or stronger] encryption). Each user must have a unique username and secret password to authenticate into the system’s UI using a browser. All passwords, whether used to access the web UI or to access a server directly via the backend, are required to meet minimum requirements for strength, complexity, and aging. Subsequent responses by the server to requests sent from the user’s browser are restricted and processed according to the user’s assigned role (i.e., using role-based access control that is strictly enforced by the application logic).

Direct access to the VM which hosts the system is only possible via a virtual private network (VPN) and then only by senior technology staff who are documented on an access control list that is reviewed for accuracy at least quarterly. Direct access to the MySQL database is also only possible by using the VPN, and the MySQL database server is configured to disallow network connections and only to accept connections that originate from the same server (“localhost”). That is, it is not possible for a MySQL client to connect to the MySQL database over the network. The client must be on the same VM as the database server.
The firewalls that implement the VPN only allow connections to ports that are required for the system to function (i.e., to port 443 for HTTPS connections). All other attempted connections are rejected. This configuration ensures that an end-user can only access data stored in the database by first authenticating with and sending requests through the web server, which processes each request and returns results that are determined by the user’s role.

The system leverages industry-standard techniques to maintain high-levels of data security and integrity. User credentials are stored as salted hashes, and the study team does not have access to or the ability to recreate user passwords. Direct access to the study database is tightly controlled. Moreover, all changes to the database (create, modify, delete) are tracked using a 21 CFR § 11-compliant audit trail.

Security is further ensured by the use of an Alert Logic Intrusion Detection System (IDS) and an Alert Logic Log Manager, both of which monitor all servers on the network. The Alert Logic IDS inspects all inbound and outbound network traffic (i.e., data packets) that passes through the network (both from the Internet and within the private network). Moreover, the IDS generates regular reports of possible exploits, thereby ensuring that outdated server software is promptly detected and upgraded to more secure patch levels.

To ensure high-availability, the system is configured with a primary server and an identical failover (“disaster recovery”) server that is located at a geographically remote data center. If the primary site becomes unavailable, the failover site can be activated well below the 60-minute Recovery Time Objective. The primary and failover database servers are also configured for replication, thereby ensuring that any transaction on the primary database also occurs on the failover database seconds later. As a further safeguard against data loss, the failover server exports the database to file every thirty minutes, and retains exports for 72 hours. In addition to the database exports, each server itself is backed up in-full daily and weekly.

To ensure that the protected health information (PHI) of research participants remains confidential, any personally identifying information (PII) that is stored in the database is encrypted using 128-bit AES encryption algorithm. Moreover, the technology staff who maintain the systems are required to use personal computers that use password protected access, screen locking, hard drive encryption, VPNs, and virus scanning utilities to ensure that their access to study’s servers cannot be compromised or misappropriated by unauthorized parties.

The data centers are monitored by certified network technicians and security personnel 24/7/365, and provide physical and logical security through a combination of keycard protocols, biometric scanning protocols, and around-the-clock interior and exterior surveillance. Access to the facilities is limited to authorized data center personnel—no one can enter the production area without prior clearance and appropriate escort. Every data center employee undergoes thorough background security checks. Conditioned power provides 100% availability through UPS (Uninterruptible Power Supply) for all servers; N+1 redundant UPS power subsystem, with instantaneous failover if the primary UPS fails. If an
extended utility power outage occurs, routinely tested, onsite diesel generators can run indefinitely. The facilities also offer a precision environment. Using an N+1 redundant HVAC (Heating Ventilation Air Conditioning) system ensures a duplicate system immediately comes online in the event of an HVAC system failure; Every 90 seconds, all air is circulated and filtered to remove dust and contaminants; the facilities also use advanced fire suppression systems appropriate for a data hosting environment. The facilities network topology and configuration was co-developed with Cisco and guards against single points of failure at the shared network level. Cisco and Arbor Networks work with the data centers to continually improve monitoring and security.

III.A.4.b. PopMedNet Data Security
Data will be transferred from the sites to the DCRI using PopMedNet. PopMedNet consists of two layers: a security layer where access controls and permissions are established and an exchange layer through which questions and responses are passed. Each of the current implementations of PopMedNet™ is hosted in a Federal Information Security Management Act (FISMA)-compliant private cloud tier III data center. All communications from the portal are encrypted. Selected current and in-process security features include:

- Strong passwords expire every 6 months and may not be reused
- Automatic logoff after inactivity
- Automated query results deletion
- Audit of all system activity
- DataMart administrators notified of new users
- Encrypted password storage
- Use of cryptographically secure random values for session IDs
- Secure distribution of DataMart Client software

III.A.4.c. Duke Clinical Research Institute Data Security
One major risk in patient-centered research is introduced by the necessity to use patient identifiers (e.g., a patient’s name and phone number is needed in order to call them for follow-up) in data systems outside the enrolling site without compromising the confidentiality and security of data. The principles of minimum-necessary and compartmentalization of identifiable data will be adopted. Primary identifiers will be retained only within the data systems where required to conduct the study; these will be omitted as data travel downstream and only study IDs will be in the analysis datasets. Further de-identification and anonymization methods will be used when making data available for secondary analysis.

The DCRI, as part of the Duke University Health System HIPAA-covered entity, routinely operates at the intersection of handling PHI and the requirements of large-scale, multicenter research programs and data-sharing initiatives. The primary computing platforms for enterprise systems are Sun Servers running the Solaris 8 operating system, a Unix-based operating system. All Oracle databases run on the Sun Servers, which are additionally attached to the HP Storage Works Storage Area Network (SAN). Client-server applications require network authentication. In addition, client-server applications may have their own internal security system and/or Relational Database Management System (RDBMS)
security. The DCRI’s core back-end tools used on this project will be SAS Analytics (SAS Institute Inc.) for data integration and data analysis. As a HIPAA-covered entity and experienced research organization, we employ a variety of technical and SOP-driven approaches to ensure the security and confidentiality of all data.

Data will be transferred from the sites to the DCRI using PopMedNet. The DCRI will be responsible for maintaining the confidentiality and security of transferred data. The control of access to databases will be managed centrally by the DCRI through user passwords linked to appropriate access privileges. This protects data from unauthorized view and modifications as well as inadvertent loss or damage. The DCRI, and our technology partner Lincoln Peak Partners, have an extensive data security infrastructure. Database servers are secured by a firewall as well as through controlled physical access. Oracle has many security protection features that ensure that each person accessing the database has the proper authority to perform the functions he or she requests of the data management system. Within the secondary SAS databases, UNIX group access control will be used for maintaining similar security. The Sun workstation log-in will be secured by extensive user password facilities under UNIX.

DCRI is part of the Duke HIPAA covered entity and has experience working with PHI in the research context; it is extremely prudent in keeping patient data secure and confidential.

III.A.5. Endpoints and Adverse Events

III.A.5.a. Primary Endpoint
The primary effectiveness analysis will be performed on the entire randomized (intention-to-treat) population. The primary endpoint of this study is the composite rate of all-cause mortality, hospitalization for nonfatal MI, or hospitalization for nonfatal stroke. Traditional reporting of potential endpoints by study sites with independent adjudication by a Clinical Events Committee will not be done in this trial given the pragmatic nature of the ADAPTABLE study and the relatively low-cost of the trial budget. We are planning to implement an endpoint validation plan to ensure the accuracy of endpoint classification compared with clinical endpoint adjudication processes used in traditional clinical trials. Once approved, the description of the validation plan will be provided in a separate document. Validated coding algorithms will be applied to a variety of EHR data sources to comprehensively ascertain potential endpoints related to hospitalizations for the non-fatal component primary endpoints (MI and stroke) and the secondary endpoints. A recently published study demonstrated that events ascertainment and classification using coding algorithms with administrative claims data yielded similar results compared with the standard adjudication of events done in traditional clinical trials. The informed consent form will cover access to the data needed to support the validation plan.

III.A.5.b. Secondary Endpoints
Secondary endpoints include the components of the primary endpoint, coronary revascularization procedures (PCI and CABG), and quality of life and functional status.
III.A.5.c. Safety
The major safety endpoint is hospitalization for major bleeding with an associated blood product transfusion.

III.A.5.d. Identifying Endpoints
Endpoints will be ascertained by applying algorithms designed to ensure comprehensive surveillance for all potential endpoints.

- Routine queries will be applied to the PCORnet CDM to capture and classify endpoints using validated coding algorithms. Because the EHR and (by extension) the PCORnet CDM is the source of truth for the trial, no patient confirmation or other additional confirmation will be required.

- Hospitalization reported by patients captured and classified by the CDM will be evaluated in the following manner:
  - Queries of the PCORnet CDM will be used to classify in-network hospitalizations (nonfatal MI, nonfatal stroke, or major bleeding).
  - It may be possible for Out-of-network hospitalizations not captured in the CDM to be investigated by executing queries against the following:
    1. Near real-time Medicare fee-for-service claims that are updated quarterly
    2. Data held by large national health plans participating in the FDA’s Mini-Sentinel initiative
    3. In the event that it is not possible to identify the patient reported event in either of the above 2 datasets, then hospitalization records will be obtained by the DCRI Call Center and used to classify the patient reported endpoints

- Death events are often not well represented in the EHR, especially out-of-hospital deaths. When participants do not respond to regular attempts at contact and no death has been reported in the PCORnet CDM, patient finder services will be utilized to find the patient. Additionally a series of cross-checks will be done to comprehensively capture all deaths that occur during trial follow-up. These cross checks for death ascertainment could be done via the Medicare Beneficiary Summary File (which includes death dates provided by the Social Security Administration) and the National Death Index.

III.A.5.d.i. Death
This endpoint includes death from any cause (all-cause mortality).

III.A.5.d.ii. Hospitalization for Nonfatal MI
The endpoint of hospitalization for nonfatal MI will be ascertained using ICD-9-CM diagnosis codes 410.x0-410.x1 in the principal or primary position. The algorithm was developed for use in FDA’s Mini-Sentinel program and has been shown to have a positive predictive value (PPV) of 86%. If these diagnosis codes are not found for a particular patient, then self-reported data regarding occurrence of nonfatal MI will be used.
III.A.5.d.iii. Hospitalization for Stroke
The endpoint of hospitalization for nonfatal ischemic stroke will be ascertained using ICD-9-CM diagnosis codes 430.x, 431.x, 433.x1, 434.x1, 435.x, 436, and 362.3 (PPV=90%). The endpoint of nonfatal hemorrhagic stroke will be ascertained using ICD-9 diagnosis codes 431 and 432 (PPV=91%). Both ischemic and hemorrhagic stroke hospitalizations will be incorporated into a combined endpoint of all stroke events. If these are not found for a particular patient, then self-reported regarding occurrence of nonfatal stroke data will be used.

III.A.5.d.iv. Coronary Revascularization
Coronary revascularization includes all coronary revascularization procedures (PCI/CABG) performed during the study. These will be identified using ICD-9-CM procedure codes (00.66, 36.06, 36.07, 00.40-00.48, 36.10-36.19) and CPT procedure codes (92920, 92921, 92924, 92925, 92928, 92929, 92933, 92934, 92937, 92938, 92941, 92943, 92944).

III.A.5.d.v. Hospitalization for Major Bleeding
Major bleeding at any location will be ascertained using previously validated ICD-9-CM diagnosis codes for a) intracranial bleeding, b) gastrointestinal bleeding, c) bleeding at another location or physician service code for GI hemorrhage (CPT code 43255 or ICD-9 procedure code 44.4x), together with a CPT code 36430 for any blood product transfusion during the same hospitalization.102

III.A.5.d.vi. ICD-10 Coding Algorithms
Given the recent implementation of ICD-10 in routine clinical practice in the United States, the aforementioned coding algorithms will be updated and adapted to ICD-10 specifications.

III.A.5.d.vii. Quality of Life and Functional Status
In addition to the effectiveness and safety endpoints listed above, we will collect data on quality of life and functional status as shown in Table 5:

<table>
<thead>
<tr>
<th>Domain</th>
<th>Item text</th>
<th>Answer list</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Health</td>
<td>In general, would you say your health is</td>
<td>5=Excellent; 4=Very good; 3=Good; 2=Fair; 1=Poor</td>
</tr>
<tr>
<td>Quality of life</td>
<td>In general, would you say your quality of life is</td>
<td>5=Excellent; 4=Very good; 3=Good; 2=Fair; 1=Poor</td>
</tr>
<tr>
<td>Physical Function</td>
<td>Are you able to run errands and shop?</td>
<td>5=Without any difficulty; 4=With a little difficulty; 3=With some difficulty; 2=With much difficulty; 1=Unable to do</td>
</tr>
<tr>
<td>Depression</td>
<td>In the past 7 days…I felt depressed</td>
<td>1=Never; 2=Rarely; 3=Sometimes; 4=Often; 5=Always</td>
</tr>
<tr>
<td>Fatigue</td>
<td>During the past 7 days…I feel fatigued</td>
<td>1=Not at all; 2=A little bit; 3=Somewhat; 4=Quite a bit; 5=Very much</td>
</tr>
<tr>
<td>Sleep Disturbance</td>
<td>In the past 7 days…I had a problem with my sleep</td>
<td>1=Not at all; 2=A little bit; 3=Somewhat; 4=Quite a bit; 5=Very much</td>
</tr>
<tr>
<td>Social Roles &amp; Activities</td>
<td>I have trouble doing all of my regular leisure activities with others</td>
<td>5=Never; 4=Rarely; 3=Sometimes; 2=Usually; 1=Always</td>
</tr>
</tbody>
</table>

III.A.5.e. Events Collection
(Refer to section III.A.5.d. Identifying Endpoints.)
These data will be analyzed, and the trial statistician will provide them to the DSMB regularly. The DSMB will review the data every 6 months and recommend any necessary changes to the conduct of the trial. Given that aspirin has been used for more than a century and most serious adverse events have been reported in conjunction with its use, the rate of unsuspected serious adverse events that the sites will/can submit through MedWatch to the FDA is expected to be very low.

III.A.6. Summary of Statistical Methods

Descriptive summaries of baseline demographic and clinical variables will be generated for each randomized treatment arm of the study. Continuous baseline variables will be presented as medians with 25th and 75th percentiles, and discrete variables will be summarized using frequencies and percentages.

III.A.6.b. Populations for Analysis
The Intent-To-Treat (ITT) Population will consist of all patients randomized to a treatment group in the study regardless of their compliance with the study medication. For all analyzed using the ITT population, subjects will be analyzed as randomized.

The per-protocol population is a subset of the ITT population excluding subjects who complied with the randomized treatment for less than 50% of the follow-up or had major protocol deviations expected to affect the primary effectiveness or safety endpoint.

III.A.6.c. Primary Effectiveness Comparison
The primary endpoint of this study will be survival free from the first event of a composite of all-cause death, hospitalization for nonfatal myocardial infarction, or hospitalization for nonfatal stroke. Specifications for the identification of these endpoints are provided in Section III.A.5. The primary effectiveness analysis will be performed by the ITT principle based on randomized treatment assignment. Event-free survival rates will be compared using Cox proportional-hazards models. In the absence of any other covariates, this is the same as the log-rank test. The test will be two-tailed and will be performed at an overall $\alpha$ of 0.05.

III.A.6.d. Primary Safety Comparison
The primary safety endpoint of this study will be the first occurrence of hospitalization for major bleeding as defined in Section III.A.5. The primary safety analysis will be performed by the ITT principle based on randomized treatment assignment. Event-free survival rates will be compared using Cox proportional-hazards models, equivalent to the log-rank test. The test will be two-tailed and will be performed at an overall $\alpha$ of 0.05.
III.A.6.e. Power

The following calculations were performed using PASS software. For the primary effectiveness endpoint, power calculations were based on an estimated primary event rate of 5% per year (in the higher dose arm), annualized rate of loss to follow-up of 5%, two-sided significance level $\alpha$ of 0.05, 10,000 patients in each treatment arm, enrollment of 24 months and a maximum follow-up period of 30 months. The power of the chosen testing strategy to detect a statistically significant difference under these assumptions is 85% if the relative risk reduction is 15%, corresponding to a total of 1308 primary effectiveness events. The power levels for other combinations of event rates and relative risk reductions, keeping all other assumptions the same, are presented in Table 6.

For the primary safety endpoint, power calculations were based on an estimated primary event rate of 2.5% per year (in the higher dose arm), annualized rate of loss to follow-up of 5%, two-sided significance level $\alpha$ of 0.05, 10,000 patients in each treatment arm, enrollment of 24 months and a maximum follow-up period of 30 months. The power of the chosen testing strategy to detect a statistically significant difference under these assumptions is 81% if the relative risk reduction is 20%, corresponding to a total of 642 primary safety events.

The power levels for other combinations of event rates and relative risk reductions, keeping all other assumptions the same, are presented in Table 7.

III.A.6.f. Secondary Endpoints

Secondary endpoints include the components of the primary endpoint, coronary revascularization procedures, quality of life and functional status (Section III.A.5.a). Time-to-event outcomes will be analyzed using the same approach as the one used for the primary endpoint. Variables collected on numerical scales will be analyzed as continuous variables. Linear repeated measures mixed model will be employed to compare the two treatments on the changes from baseline.
III.A.6.g. Prior Treatment Effect
We expect that most patients recruited into the ADAPTABLE trial will already be treated with pre-randomization aspirin therapy, given the nature of the inclusion criteria and the known high utilization of aspirin for the secondary prevention of coronary artery disease in the United States. Therefore, a sensitivity analysis with a 10-day landmark will be performed that excludes all events occurring in the first 10 days following randomization in order to formally test for a legacy effect of aspirin.

III.A.6.h. Handling of Missing Data
Concerted effort will be made to eliminate or minimize the occurrence of missing data. Participants will grant access to their electronic medical records at enrollment and during study follow-up as well as to have their information searched in national databases. If, despite these efforts, missing data occur, we will employ the following statistical techniques to address them.

First, reasons for missing data will be collected and described. All patients will be accounted for in all analyses and presentations. For the primary analysis based on event-free survival, subjects discontinuing the study prematurely will be censored at the time of discontinuation. However, this approach might lead to biased results if discontinuation does not occur at random. Thus, two “sensitivity analyses” will be undertaken:

1. **Inverse probability weighting.** In this approach, the contribution of each subject to the risk set calculated at time $t$ will be inversely weighted by the estimated probability of remaining uncensored up to time $t$. This probability will be estimated using a Cox proportional hazards model fitted to time to censoring with variables potentially prognostic of both, failure and censoring, with baseline and time-dependent (such as most frequent major protocol deviations, certain AEs etc.), entered as covariates. In order to reduce the potentially high variability of the resulting treatment effect estimators due to sampling variability in weights, the weights will be “stabilized” by multiplication of probabilities of remaining uncensored up to time $t$ estimates using baseline covariates only.

2. **Pattern-mixture approach.** Following Little et al.\textsuperscript{106,107} we will assume that for participants who drop out, the hazard of an outcome deviates from that of participants who do not drop out by an offset, denoted by $d_1$ for the higher dose and by $d_0$ for the lower dose. We will then explore the effect of this deviation on the findings for various choices of the offsets in the two study groups. If the treatment effect is qualitatively maintained for the range of offsets that are considered to be clinically plausible, then the findings will be considered to be robust.

In other analyses, missing data will be handled using multiple imputations. Ten imputed data sets will be generated with imputation methods based on the regression or Monte Carlo framework. Final results will be based on averages from the 10 imputed data sets with appropriate estimator employed of the variance.\textsuperscript{107}
III.A.6.i. Subgroup Analyses (Heterogeneity of Treatment Effect)

Subgroup analyses for the primary effectiveness and safety endpoints will be performed on the ITT population in order to explore whether the treatment effect is consistent across subgroups. Subgroup analyses to evaluate variation in treatment effect will be performed on the basis of tests for interaction using the Cox proportional hazards model with terms for treatment group, the subgroup variable and treatment by subgroup variable interaction. Additionally, treatment effects within each categorical subgroup will be examined separately using Cox proportional hazards models. Event rates by treatment and HRs with 95% confidence intervals will be reported for each subgroup. Forest plots will be generated displaying the estimated hazard ratios and 95% confidence intervals for each subgroup will be presented. For subgroups defined using continuous variables, the analysis based on the continuous form will be considered primary but for display purposes these variables can also be categorized.

The following subgroups determined at baseline will be examined:

- Age ≥ 65 years
- Race categories (White, Black, and Asian; Hispanic ethnicity)
- Diabetes mellitus
- Chronic kidney disease (serum creatinine > 1.5 mg/dL)
- Current P2Y12 inhibitor use
- Female sex

Although the importance of understanding the effects of aspirin in these subgroups is critical, we recognize that despite enrolling over 20,000 participants, our power to detect statistically significant and clinically meaningful interactions will be limited. However, this effort represents the largest such undertaking and is of vital importance. Heterogeneity of treatment effect will be established based on the interaction test specified above. Testing for differences between treatment arms within subgroups will be considered exploratory and no claims of heterogeneity will be made based on tests within subgroups. By their nature, these tests have low power unless the effect sizes are large, as illustrated in Table 8.

Patients without internet access will not be analyzed as a pre-specified sub-group, but sensitivity analyses for the main trial endpoints will be incorporated into the statistical analysis plan to analyze the populations with vs. without Internet access to assess the consistency of the treatment results and potential treatment interactions, given expected demographic and socioeconomic differences between those with vs. without internet access.

<table>
<thead>
<tr>
<th>Subgroup size as % of total</th>
<th>Relative risk reduction</th>
<th>Power</th>
</tr>
</thead>
<tbody>
<tr>
<td>10%</td>
<td>25%</td>
<td>37%</td>
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<tr>
<td>20%</td>
<td>25%</td>
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<tr>
<td>15%</td>
<td>16%</td>
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<tr>
<td>30%</td>
<td>25%</td>
<td>81%</td>
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<tr>
<td>20%</td>
<td>60%</td>
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<td>15%</td>
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<td></td>
</tr>
<tr>
<td>15%</td>
<td>81%</td>
<td></td>
</tr>
</tbody>
</table>

III.A.6.j. Interim Blinded Trial Monitoring

During the conduct of the trial, we will employ state of the art statistical monitoring techniques. These will be conducted in a blinded fashion. We will observe study discontinuation patterns as well as reasons for the missing data. We will monitor the event rate as it accumulates and develop event rate projections which will help us determine if and when we are likely to achieve sufficient power. We will
also review the baseline characteristics of enrolled participants to make sure we are enrolling the population defined in the protocol. We will also monitor the primary safety events as they accumulate.

III.A.7. Study Coordination and Monitoring

III.A.7.a. Site Management and Quality Assurance

This trial will be monitored using quality-by-design principles and will not require on-site monitoring.

III.A.7.a.i. ADAPTABLE Quality-by-Design

Final success of any clinical trial in answering the question of interest depends in large part on its design. The term “quality by design” refers to steps taken at the trial design stage to foresee and limit problems that might occur during the trial conduct. Adherence to the following five “guiding principles” greatly increases the chances of final success:

1. Have we enrolled the right participants according to the protocol with adequate consent (Right Patient)?
2. Did participants receive the assigned treatment and did they stay on the treatment (Right Treatment)?
3. Was there complete ascertainment of primary and secondary outcome data (Right Outcomes)?
4. Was there complete ascertainment of primary and secondary safety data (Right Outcomes)?
5. Were there any major Good Clinical Practice (GCP)-related issues?

This trial was designed to maximize the likelihood that the five principles noted above will be followed. In particular, patient enrollment and consent is facilitated by the carefully selected and operationalized CDRNs that are the cornerstone of the PCORnet initiative. Patient selection facilitated by electronic health records (EHRs) should lead to efficient identification and enrollment of patients who satisfy study eligibility and enrichment criteria. These criteria have been chosen in a manner that corresponds to what is routinely available in EHRs, reducing the risk of any ambiguity and enrollment of patients who do not meet the protocol-specified criteria.

Although it is not possible to guarantee that all study participants will remain on the trial until its end and fully adhere to the study drug, the existence of the ADAPTABLE patient portal and potential follow-up from the DCRI Call Center are expected to keep participants engaged and help them maintain their assigned treatment by regularly asking them about the treatment they take. At the same time, because this trial is intended to describe what happens in a “real-world” population (intent-to-treat principle), variability in treatment adherence may actually contribute to better generalizability of study results.

Similarly to the eligibility criteria, primary efficacy and safety endpoints have been defined in a way that is well synchronized with the endpoint collection tools that will be employed. The focus on “hard” endpoints (that is, endpoints that can be unambiguously determined), cause-specific hospitalizations for MI, stroke, and bleeding, greatly reduces the potential of recall bias by study participants during self-report. At the same time, it facilitates improved ascertainment through the corresponding EHRs. Accordingly, the case report forms are short and questions are phrased in a manner that does not overwhelm study participants. Access to EHRs and consent to search death records should also reduce the amount of missing primary endpoint data. Minimal levels of intervention and a focus on observing
rather than influencing the study participants greatly increases the likelihood that Good Clinical Practices will be followed.

**III.A.7.b. Study Network**

The Study Network is composed of most of the CDRNs and 1 PPRN within PCORnet. Each of these CDRNs have developed a network specific plan to engage clinicians, identify and enroll participants and be available as patients are followed. Each CDRN will strive to be activated to begin enrollment during the first year of the study. The Coordinating Center will perform specific centralized functions; this will be advantageous to meet study budget limitations.

**III.8.b.i. ADAPTABLE Leadership Structure**
III.A.7.c. Executive Committee
The ADAPTABLE Executive Committee will provide oversight and guidance for overall trial activities. It will also serve as a communication nexus to promote sharing of information and cross-pollination of ideas among the Executive Committee, Steering Committee, study co-PIs, Coordinating Center PIs and staff, PCORI representatives, and the Co-Chairs of the Patient Representatives/Adaptors group, all of whom will have representation on the Committee. The decision-making structure and process for the Executive Committee will be described in a separate Executive Committee charter.

III.A.7.d. Steering Committee
The trial’s Steering Committee will be responsible for the protocol and the scientific conduct of the study. The Committee will include patient and physician representatives from each participating CDRN and PPRN, along with selected network members with expertise in clinical research in antiplatelet therapies, cardiovascular disease, and analytical methods. Additionally, representatives from the American College of Cardiology (ACC), the American Heart Association (AHA), and PCORI will be on the Steering Committee. The Steering Committee will make decisions about any changes to the protocol and will review all proposed substudies. The decision-making structure and process for the Steering Committee will be described in a separate Steering Committee charter.

III.A.7.e. Data and Safety Monitoring Board (DSMB)
The DSMB will consist of independent members who are not participating investigators in this trial. The committee will include a Chair who is a cardiovascular specialist, a statistician, 2 patient representatives, and 3 cardiovascular specialists. The DSMB will review data regularly and recommend necessary changes to the conduct of the trial. For example, if significantly large and important treatment differences are observed during any of the interim analyses, the DSMB may recommend that randomization of patients be stopped, or that the design and conduct of the trial be appropriately modified.

We anticipate that the DSMB will meet at approximately 6-month intervals to review the accumulating data. Before each DSMB meeting, the Data Coordinating Center will conduct the desired statistical analyses and prepare a summary report for review by the DSMB. The extracted data files and analysis programs for each DSMB report will be maintained at the Data Coordinating Center for the life of the study. Each report will describe the progress in enrollment, the rates of compliance with therapy. The DSMB may recommend stopping the trial for any safety-related concern at any time. Particular attention will be paid to all-cause mortality. For effectiveness, the trial will compare doses of aspirin that have been proven effective in previous trials and thus it is unlikely that any dramatic differences will appear early. However, the DSMB will monitor these differences about every 6 months over the 2.5 years of the trial. Thus, we anticipate four interim analyses and one final analysis. Because there are multiple tests, each test must be made at an adjusted \( \alpha \) level using group sequential-testing methods. To minimize the chance of stopping early due to a spurious result, the endpoints will be tested at the planned analyses using a specific method known as the Haybittle-Peto rule. This rule tests the endpoints at the 0.001 level (\( z = 3.29 \), two-sided) at each interim review and then makes the final test at the 0.0499 level (\( z = 1.9605 \)). Thus, there is only minimal penalty for interim analyses.
The decision-making structure and process for the DSMB will be described in a separate DSMB charter.

III.A.7.f. Data Auditing Conventions
Data will be collected via the study patient portal (Section III.A.3.d). Data will be queried at the time of data entry through the study patient portal for missingness and inconsistencies.

IV. Human Subjects

IV.A. Protection of Human Subjects

IV.A.1. Human Subjects Involvement and Characteristics

IV.A.1.a. Inclusion Criteria

1. Known atherosclerotic cardiovascular disease (ASCVD), defined by a history of prior myocardial infarction, prior coronary angiography showing ≥75% stenosis of at least one epicardial coronary vessel, or prior coronary revascularization procedures (either PCI or CABG)

2. Age ≥ 18 years

3. No known safety concerns or side effects considered to be related to aspirin, including
   a. No history of significant allergy to aspirin such as anaphylaxis, urticaria, or significant gastrointestinal intolerances
   b. No history of significant GI bleed within the past 12 months
   c. Significant bleeding disorders that preclude the use of aspirin

4. Access to the Internet. In the event that the CDRNs are notified that a cohort of patients without internet access can be included, then patient agreement will be obtained during the consent process to provide follow-up information by telephone contact with the DCRI Call Center.

5. Not currently treated with an oral anticoagulant – either warfarin or a novel anticoagulant (dabigatran, rivaroxaban, apixaban, edoxaban) – and not planned to be treated in the future with an oral anticoagulant for existing indications such as atrial fibrillation, deep venous thrombosis, or pulmonary embolism.

6. Not currently treated with ticagrelor and not planned to be treated in the future with ticagrelor.

7. Female patients who are not pregnant or nursing an infant

8. Estimated risk of a major cardiovascular event (MACE) > 8% over next 3 years as defined by the presence of at least one or more of the following enrichment factors:
   a. Age > 65 years
   b. Serum creatinine > 1.5 mg/dL
   c. Diabetes mellitus (Type 1 or Type 2)
   d. 3-vessel coronary artery disease
   e. Cerebrovascular disease and/or peripheral arterial disease
   f. Left ventricular ejection fraction (LVEF) < 50%
g. Current cigarette smoker.

There will be no exclusions for any upper age limit, comorbid conditions, or concomitant medications other than oral anticoagulants and ticagrelor that are used at the time of randomization, or are planned to be used during the study follow-up.

Simple, inclusive eligibility criteria will make enrollment easier, and will render study results more generalizable to a broader population of patients. We will exclude pregnant or lactating women (because of concern for the fetus or child), patients taking oral anticoagulants or likely to require an oral anticoagulant during trial follow-up (because of complex drug interactions and a projected excessive risk of bleeding), and patients at relatively low risk for cardiovascular events (ie, no enrichment factor because of the large number of outcomes needed to detect a clinically meaningful difference with the available sample size).

IV.A.1.b. Sources of Material

Data obtained from the study patient portal and from the CDM will be transferred to the DCRI along with unique patient identifiers. Data contained with the CDM will be obtained at each of the sites and transferred to the DCRI. The control of access to databases at DCRI will be managed centrally by the DCRI through user passwords linked to appropriate access privileges. This protects data from unauthorized view and modifications as well as inadvertent loss or damage. Within the secondary SAS databases, UNIX group access control will be used for maintaining similar security. The Sun workstation login will be secured by extensive user password facilities under UNIX.

IV.A.1.c. Potential Risks

Aspirin is approved by the United States Food and Drug Administration (FDA) for the secondary prevention of ischemic events associated with ASCVD. In patients with established atherosclerosis, the risk of aspirin is quite low compared with the benefit. A very small number of patients have a serious anaphylactic reaction or bronchoconstriction with nasal polyps when they use aspirin. The risk of intracranial hemorrhage is <0.04% per year. A modest number of patients develop serious gastrointestinal bleeding as a result of loss of the protective effect of prostaglandins on the gastric mucosa. A larger number of patients have gastrointestinal intolerance, but this is often transient and can be overcome. The absolute risk varies as a function of the trial entry criteria, but none of these major risks exceeds 5 events per 100 patients treated per year. Except for patients with preexisting asthma, no studies have described risk factors for these complications of aspirin use. However, because patients will be selected based on the absence of such known major intolerances to a dose of 325 mg of aspirin, the expected risk of these major events in this trial will be low.

IV.A.1.d. Recruitment and Informed Consent

Each CDRN will develop a recruitment process that will work best within their organization, utilizing the tools they have available and their local infrastructure. Prior to cohort identification, site investigators will be asked to endorse the protocol, and depending upon their preference, they will either determine each patient’s eligibility or will give permission for the CDRN, through its integrated health system members, to identify and contact potentially eligible patients. In the latter case, patients who meet
criteria for secondary prevention after a cardiovascular event will be identified using customized search algorithms unique to each network from their aggregated EHR systems. Broad agreement from both cardiovascular specialists and primary care physicians will be sought. Patients will be enrolled after providing electronic informed consent.

**IV.A.1.e. Protection Against Risk**

Please see Sources of Material above and Data and Safety Monitoring Plan below.

**IV.A.1.f. Potential Benefits of the Proposed Research to the Research Participants**

As discussed in Section I.A., atherosclerosis leading to thrombotic events, and in particular ASCVD, represents the leading cause of death, morbidity, and disability and affects more than 150 million people worldwide. Despite remarkable progress in preventive and interventional approaches to atherosclerosis, ASCVD is expected to be an even more prominent cause of death and disability for the next 30 years. In technologically-developed countries, the major factor that contributes to this expansion is the aging of the population, such that, despite declining age-specific disease rates, the total disease burden increases because ASCVD eventually strikes a larger population of older adults. In developing countries, a major epidemic of atherosclerosis is occurring, concentrated at younger ages and presumably due to the spread of tobacco use, Westernization of diet, and a sedentary life-style.

The development of new biological and technological approaches to treating ASCVD is exciting, but maximizing the use of an inexpensive yet effective therapy is far more promising in reducing death and disability on a global scale. Numerous clinical trials have shown the clinical benefit of aspirin versus placebo in reducing vascular events, but the best dose of aspirin for the general population with ischemic heart disease has not been determined. Considering the global burden of ASCVD and that the affected population is growing rapidly, identifying the optimal dose of aspirin would save lives and prevent ischemic and bleeding events globally. For example, based on recent evidence suggesting a reduction in ischemic events with lower doses of aspirin, the odds ratio for an event with a dose of 81 mg versus 325 mg per day would be 0.84 (95% confidence interval [CI], 0.64–1.1). If borne out in a prospective clinical trial, and if the rate of death, MI, or stroke over ~18 months of treatment was 8% with 325 mg of aspirin (based on contemporary trials of aspirin use in patients with ischemic heart disease), then the expected event rate with 81 mg would be 6.8% (95% CI, 5.3–8.7), or ~12 events prevented for every 1000 patients treated. Considering the increasing global burden of ischemic heart disease, a 1.2% absolute reduction in events simply through optimal aspirin dosing would be of tremendous importance to public health.

**IV.A.1.g. Importance of the Knowledge to Be Gained**

The primary aim of this study is to determine the optimal dose of aspirin, the universally available effective therapy for ASCVD. Despite dozens of clinical trials involving over 100,000 patients, the most effective aspirin dose has not been identified. Nonrandomized studies have suggested that lower doses of aspirin may be associated with not only lower rates of bleeding but also reduced ischemic outcomes. Despite guideline recommendations for lower dose aspirin, over half of post-ACS patients in the US are discharged on a dose of 325 mg. Considering the worldwide epidemic of ASCVD and the potential
benefits of aspirin, identifying the optimal dose of aspirin would have substantial global health and economic effects similar to few other medical therapies.

Although the primary aim of this study is to determine the optimal dose of aspirin, this study also represents a new and efficient interactive model for designing and implementing clinical trials that aim to refine therapies already in use in contemporary clinical practice. Because we live in an era in which the number of effective (or potentially effective) therapies far exceeds our ability to evaluate them in prospective clinical trials using current methods, there is an urgent need to develop an approach to outcome-based trials that can greatly reduce the cost per patient. By following patients directly on the Internet and thereby avoiding the costs of clinic visits, lengthy case record forms and extensive site management, we believe that a new, more efficient, and less expensive model for trials can be developed that could be extended to more experimental comparisons. However, working through the issues of informed consent, data validation, events ascertainment, and compliance assessment will require acceptance of novel approaches to statistical sampling and an intense focus on communication with both patients and their physicians. In this era of increasing concern about both patient privacy and research integrity, such an approach to trial efficiency would be exceedingly difficult to pilot with untested therapies. Because this trial will test only doses of aspirin that are considered relatively safe and are widely used in current clinical practice, it provides a critically and globally important clinical issue with which to develop such new clinical trials methods.

This trial also provides an opportunity to explore the use of integrated health systems, EHRs, and patient reported outcomes as tools for performing clinical trials. A trial conducted almost exclusively over the Internet offers many potential benefits. First, patients would have access to a custom-built, patient centered patient portal that offers current information about symptom awareness, risk factor modification, and disease management and prevention. The patient portal would also serve as the primary mechanism for follow-up, with routine data entry by the patients themselves. Not only would this method examine the practicability of patient self-reporting, but it would potentially enable substantial savings in cost and resources (e.g., costs associated with physician reporting). For physicians, use of the Internet in clinical trials could further broaden awareness and participation and, at the same time, facilitate the conduct of the study. For example, trial enrollment and follow-up could be immediate with use of the Internet, at any time, eliminating a need for traditional methods (e.g., face to face visits). Finally, the platform created by this trial will unite a far-reaching community of patients and their physicians with a common goal of refining an existing therapy to maximize its benefit relative to risk. It is likely that such knowledge will produce far greater global benefit than the introduction of many other “high-tech” approaches.

IV.B. Inclusion of Women
(see following section)

IV.C. Inclusion of Minorities
This Study will seek to enroll a diverse population representative of the broad population of patients with ASCVD. The CDRN medical practices are from communities representing a wide range of sex, age,
racial, and ethnic backgrounds. We will make every attempt to explain the project in easy (non-clinical) terms to all patients to make sure they understand the importance of the research and the need to have good representation of key subgroups and they appreciate the potential benefit they could derive from participating in it. A particular advantage for this trial is the relative lack of barriers to participation. Because clinic visits are not needed for follow-up, additional transportation is not required, and there is minimal additional time commitment for patients. With regard to the specific enrollment of women, this is discussed in detail in Section III.A.2.a-b. Strategies to enhance the enrollment of ethnic minorities have been described in detail in Section III.A.2.a-b. We will strive to enroll a proportion of minority patients similar to that in the overall ASCVD population. Content on the ADAPTABLE patient portal will be available in English and Spanish.

IV.D. Inclusion of Children
Because participation in this trial is restricted to patients aged ≥18 years, inclusion of children in this trial is prohibited. Since the incidence of coronary artery disease among young (i.e., <18 years) individuals is very low, children do not represent a study population relevant to this trial.

IV.E. Data and Safety Monitoring Plan
This has been described in detail in Section III.A.7 (Study Coordination and Monitoring).
Appendix: Patient Survey

Aspirin Trial Questions

Have you had a history of MI (myocardial infarction?)
- Yes
- No
- Don't know

Have you had a history of CAD (Coronary artery disease?)
- Yes
- No
- Don't know

Do you take any of these medications on a regular basis?
Mark all that apply
- Aspirin 81mg "baby aspirin"
- Aspirin 325mg "regular aspirin"
- Aspirin at another dose
- Apixaban (Eliquis)
- Clopidogrel (Plavix)
- Dabigatran (Pradaxa)
- Prasugrel (Effient)
- Rivaroxaban (Xarelto)
- Ticagrelor (Brilinta)
- Warfarin (Coumadin)
Aspirin Trial Questions

**What dose of aspirin do you take?**
Answer only if applicable

**If you take Aspirin: How often do you usually take it?**
Answer only if applicable
- Every day, 2 times per day
- Every day, 1 time per day
- Every other day
- Less often

**Have you ever had to stop taking aspirin because of side effects like a rash, upset stomach or bleeding?**
Answer only if applicable
- Yes
- No

**If you had side effects from taking aspirin please list them**
Answer only if applicable

**Would you be willing to be a patient advisor to this study?**
This may include: helping with study design, identifying ways to recruit study participants, helping develop a patient-friendly consent form, helping identify study outcomes important to patients, participating in the study's data safety monitoring process and helping interpret and disseminate results.
- Yes
- No

*Never submit passwords through Google Forms.*
References


42. Wiviott SD, Braunwald E, Angiolillo DJ, et al. Greater clinical benefit of more intensive oral antiplatelet therapy with prasugrel in patients with diabetes mellitus in the trial to assess improvement


84. Williams MA, Fleg JL, Ades PA, et al. Secondary prevention of coronary heart disease in the elderly (with emphasis on patients > or =75 years of age): an American Heart Association scientific statement


