Brazilian guidelines on prevention of cardiovascular disease in patients with diabetes: a position statement from the Brazilian Diabetes Society (SBD), the Brazilian Cardiology Society (SBC) and the Brazilian Endocrinology and Metabolism Society (SBEM)

Marcello Casaccia Bertoluci1,2*, Rodrigo Oliveira Moreira3,4,5, André Faludi6, Maria Cristina Izar7, Beatriz D. Schaan9, Cynthia Melissa Valerio3, Marcelo Chiara Bertolami9, Ana Paula Chacra9, Marcus Vinicius Bolivar Malachias10, Sérgio Vencio11, José Francisco Kerr Saraiva12, Roberto Betti9, Luiz Turatti9, Francisco Antonio Helfenstein Fonseca7, Henrique Tria Bianco7, Marta Sulzbach6, Adriana Bertolami9, João Eduardo Nunes Salles13, Alexandre Hohl14, Fábio Trujilho15, Eduardo Gomes Lima9, Marcio Hiroshi Miname9, Maria Teresa Zanella18, Rodrigo Lamounier16, João Roberto Sá19, Celso Amodeo6, Antonio Carlos Pires17 and Raul D. Santos9

Abstract

Background: Since the first position statement on diabetes and cardiovascular prevention published in 2014 by the Brazilian Diabetes Society, the current view on primary and secondary prevention in diabetes has evolved as a result of new approaches on cardiovascular risk stratification, new cholesterol lowering drugs, and new anti-hyperglycemic drugs. Importantly, a pattern of risk heterogeneity has emerged, showing that not all diabetic patients are at high or very high risk. In fact, most younger patients who have no overt cardiovascular risk factors may be more adequately classified as being at intermediate or even low cardiovascular risk. Thus, there is a need for cardiovascular risk stratification in patients with diabetes. The present panel reviews the best current evidence and proposes a practical risk-based approach on treatment for patients with diabetes.

Main body: The Brazilian Diabetes Society, the Brazilian Cardiology Society, and the Brazilian Endocrinology and Metabolism Society gathered to form an expert panel including 28 cardiologists and endocrinologists to review the best available evidence and to draft an up-to-date an evidence-based guideline with practical recommendations for risk stratification and prevention of cardiovascular disease in diabetes. The guideline includes 59 recommendations covering: (1) the impact of new anti-hyperglycemic drugs and new lipid lowering drugs on cardiovascular risk; (2) a guide to statin use, including new definitions of LDL-cholesterol and in non-HDL-cholesterol targets; (3) evaluation of silent myocardial ischemia and subclinical atherosclerosis in patients with diabetes; (4) hypertension treatment; and (5) the use of antiplatelet therapy.
Background
Since the first position statement on diabetes and cardiovascular prevention published in 2014 by the Brazilian Diabetes Society [1], important studies have been published in the area of cardiovascular assessment and prevention in patients with diabetes [2]. These studies have deeply advanced the current view on primary and secondary prevention in diabetes, and suggested new approaches on cardiovascular risk stratification, new cholesterol-lowering drugs, and new anti-hyperglycemic drugs with novel significant cardiovascular effects and mortality reduction.

To address this challenge, and in recognition of the multifaceted nature of disease, the Brazilian Diabetes Society joined the Brazilian Society of Cardiology and the Brazilian Endocrinology and Metabolism Society and gathered an expert panel formed by 28 cardiologists and endocrinologists to review the best available evidence and to draft up-to-date evidence-based guidelines with practical recommendations on both the stratification and prevention of cardiovascular disease in diabetes. The main innovations include: (1) considerations on the impact of new anti-hyperglycemic drugs and new lipid-lowering drugs on cardiovascular risk; (2) a practical risk factor-based approach to guide statin use, including new definitions of LDL-cholesterol and non-HDL-cholesterol targets; (3) an evidence-based approach to evaluate silent myocardial ischemia and subclinical atherosclerosis in patients with diabetes; (4) the best current approaches for treating hypertension; and (5) recommendation updates for the use of antiplatelet therapy. We hope these guidelines will help clinicians to improve the quality of the care provided to patients with diabetes.

Methods
Initially, the panel members were divided into seven subcommittees to define the main topics requiring an updated position from the societies. Panel members searched PUBMED for randomized clinical trials and meta-analyses of clinical trials, and observational studies of good quality published from 1997 to 2017 using MeSH terms: [diabetes], [type 2 diabetes], [cardiovascular disease], [cardiovascular risk stratification] [coronary artery disease], [screening], [silent ischemia], [statins], [hypertension], [acetyl salicylic acid]. Low quality observational studies, meta-analyses with high heterogeneity and cross-sectional studies were not included although they might have influenced the level of evidence indicated. Expert opinion was used when the results of the search were not satisfactory for a specific item. It is important to note that it was not the aim of this position statement to include a rigorous systematic review.

A preliminary manuscript outlining recommendation grades and levels of evidence (Table 1) was then drafted. This step took several rounds of discussion among subcommittee members, who reviewed the findings and made new suggestions. The manuscript was

Table 1 Recommendation grades and levels of evidence

| Grade of recommendation | Evidence is conclusive or, if not, there is a general consensus that a procedure or a treatment is safe and efficacious |
|-------------------------|---------------------------------------------------------------------------------------------------------------|
| Class I                 | There is conflicting evidence or divergent opinion on safety, efficacy or utility of treatment or procedure |
| Class II                | Opinions are in favor of the treatment or procedure. The majority of experts approves |
| Class IIa               | Less well established efficacy, opinions are divergent |
| Class IIb               | There is evidence or consensus that the treatment or procedure is not useful, efficacious or may be harmful |

| Levels of evidence | Multiple concordant well designed randomized clinical trials or robust meta-analyses of randomized clinical trials |
|--------------------|---------------------------------------------------------------------------------------------------------------|
| A                  | Data from less robust meta-analyses, a single randomized clinical trial or observational studies |
| B                  | Expert opinion |

Conclusions: Diabetes is a heterogeneous disease. Although cardiovascular risk is increased in most patients, those without risk factors or evidence of sub-clinical atherosclerosis are at a lower risk. Optimal management must rely on an approach that will cover both cardiovascular disease prevention in individuals in the highest risk as well as protection from overtreatment in those at lower risk. Thus, cardiovascular prevention strategies should be individualized according to cardiovascular risk while intensification of treatment should focus on those at higher risk.

Keywords: Diabetes mellitus, Cardiovascular prevention, Cardiovascular screening, Blood glucose, Risk factors, Coronary artery disease, Dyslipidemias, Hypertension, Antiplatelet agents
then returned to the lead author in charge of text harmonization and inclusion of minor changes, and was subsequently submitted to further view rounds by committee members, seeking a consensus position. After this phase, the manuscript was forwarded to the editorial board for final editing and submitting for publication.

These guidelines were divided into seven modules, namely:

**Cardiovascular risk**
- Module 1: Cardiovascular risk stratification
- Module 2: Screening of subclinical atherosclerosis
- Module 3: Screening of silent myocardial ischemia

**Cardiovascular prevention**
- Module 4: Management of hyperglycemia
- Module 5: Management of dyslipidemia
- Module 6: Management of hypertension
- Module 7: Antiplatelet therapy

### Module 1: Cardiovascular risk stratification

Patients with type 1 and 2 diabetes are divided into four broad cardiovascular risk categories—LOW, INTERMEDIATE, HIGH, AND VERY HIGH (Table 2)—based on age, presence of stratifying risk factors (SF) (Table 3), subclinical atherosclerosis (SCAT) (Table 4), or clinical atherosclerotic disease (CLAD) (Table 5). The 10-year cardiovascular event rate for low, intermediate, high, and very high risk categories were respectively: <10, 10–20, 20–30, and >30% (Table 2).

The LOW and INTERMEDIATE risk categories are based solely on age and SF (Table 3). SCAT (Table 4), and CLAD (Table 5) are not present in these risk groups. As seen in a large Ontario population-based retrospective cohort study, 379,003 individuals with diabetes were included and followed up for a mean of 8 years until the occurrence of a first acute myocardial infarction or death from all causes [3]. The transition from LOW to INTERMEDIATE RISK occurred at ages 38 and 46 years respectively for men and women. The transition from INTERMEDIATE to HIGH-RISK status occurred

### Table 2  Cardiovascular risk categories in patients with diabetes

| Risk category | CHD event rate in 10 years (%) | Age | Condition |
|---------------|---------------------------------|-----|-----------|
| LOW | <10 | Men < 38 years  Women < 46 years | No stratification factors (SF)<sup>a</sup> |
| INTERMEDIATE | 10–20 | Men 38–49 years  Women 46–56 years | No subclinical atherosclerosis (SCAT)<sup>b</sup> |
| HIGH | 20–30 | Men > 49 years  Women 56 years or any age if SF<sup>a</sup> or SCAT<sup>b</sup> | Stratification factors (SF)<sup>a</sup> |
| VERY HIGH | >30 | Any age if CLAD<sup>c</sup> | Clinical atherosclerotic disease (CLAD)<sup>c</sup> |

<sup>a</sup> Stratification factors (Table 3)
<sup>b</sup> Subclinical atherosclerosis (Table 4)
<sup>c</sup> Clinical atherosclerotic disease (Table 5)
respectively at ages: 49 and 56, for both men and women [3]. Therefore, patients with diabetes without clinical or subclinical cardiovascular disease and risk factors are considered at INTERMEDIATE RISK when aged are 38–49 years (men) or 46–56 years (women), and at LOW RISK if they are younger.

The HIGH-RISK group is defined by the presence, at any age, of at least one SF (Table 3) or one indicator of SCAT (Table 4), in the absence of CLAD (Table 5). Even in the absence of these conditions, a patient with diabetes is also considered at HIGH RISK when age is above 49 years in men or 56 years in women. Finally, the VERY HIGH-RISK group includes patients who, at any age, have CLAD as defined in Table 5.

Module 2: Screening of subclinical atherosclerosis

1. Coronary artery calcification (CAC) score is associated with cardiovascular events and mortality in patients with diabetes [I, A]

Summary of evidence

- Coronary artery calcification (CAC) is a marker for the presence and burden of atherosclerosis, as demonstrated in anatomical studies [22]. The MESA [23] and Heinz Nixdorf [24] studies demonstrated that CAC is a predictor of coronary events and is useful for stratification of cardiovascular risk among patients in primary prevention. This is also true for patients with diabetes: the higher the CAC score, the higher the risk of cardiovascular events in subjects with diabetes [25].

- Raggi et al. [26] followed 10,377 asymptomatic individuals (903 with diabetes), who had been investigated with CAC at baseline, for a mean of 5 years. The mean CAC score was higher in patients with diabetes than in patients without diabetes (281 ± 567 vs. 119 ± 341, p < 0.0001). This study also showed that a higher CAC score was associated with a higher mortality rate, especially in patients with diabetes. However, the survival rate was similar in patients with and without diabetes (98.8% vs. 99.4% respectively, p = 0.5) when CAC was zero.

- The PREDICT study [27] followed 589 patients with diabetes without cardiovascular disease (mean age 63.1 years) for a median of 4 years. The greater the coronary calcium score, the greater the risk of cardiovascular outcomes. The area under the ROC curve (AUC-ROC) for risk determination using the UKPDS risk score was 0.63, and was increased to 0.73 when CAC was included (p = 0.03).

2. Coronary artery calcium score (CAC score) determination has the best net reclassification rate compared to other risk markers when added to clinical global risk score calculators alone. This can be especially useful to reclassify patients at INTERMEDIATE risk to higher or lower-risk categories. However, this Panel recognizes that, despite its utility, CAC score may not be easy to obtain in a large proportion of patients [IIa, B]

Summary of evidence

- In a large cohort study of 44,052 asymptomatic individuals referred for CAC testing, including 2384 with diabetes [28], the authors showed that cardiovascular risk was more accurately stratified with CAC in patients with diabetes. Patients in the low and moderate risk categories had a mortality rate of 39.4 deaths/1000/year when CAC was above 100. Conversely, those classified in the clinical high-risk category with no calcium present (CAC = 0) had a 10-year mortality rate of 6.59 deaths/1000/year. In the lower-risk subgroup (<5% in 10 years), 18% had CAC > 100, while in the higher risk category (>20% in 10 years), 16% had CAC = 0. In other words, CAC was able to reclassify a considerable number of low-risk patients into a high-risk category [27]. A CAC score >0 was present in 57.3% of patients in the low-risk category and in 70.6% of those in intermediate-risk categories.

- The prospective, community-based coronary artery risk development in young adults (CARDIA) study [29] recruited 5115 participants aged 18–30 years, with CAC measured at 15, 20, and 25 years after recruitment. The main outcomes were incident coronary heart disease, including fatal and nonfatal myo-

### Table 4 Subclinical atherosclerosis (SCAT)

| Condition                                      |
|------------------------------------------------|
| Coronary artery calcium score (CAC) >10 U Agatston⁴ |
| Carotid plaque (intima-media thickness >1.5 mm) ¹⁴  |
| Computed tomography coronary angiography (CCTA) with a definite plaque ¹⁵  |
| Ankle-brachial index <0.9 ¹⁶                     |
| Abdominal aortic aneurysm (AAA)  [17–21]²⁶      |

⁴ When available, CAC scoring should be the preferred modality

¹⁵ CCTA should not be performed routinely in truly asymptomatic patients

²⁶ Patients suffering from an AAA are at elevated risk of cardiovascular morbidity and mortality, due to common risk factors and comorbidities associated with the aneurysm

### Table 5 Clinical atherosclerotic disease (CLAD)

| Condition                                      |
|------------------------------------------------|
| Acute coronary syndrome:                       |
| Acute myocardial infarction or unstable angina |
| Stable angina or previous acute myocardial infarction |
| Atherothrombotic stroke or transient ischemic attack |
| Coronary, carotid, or peripheral revascularization |
| Peripheral vascular insufficiency or limb amputation |
| Severe atherosclerotic disease (stenosis >50%) in any vascular territory |

Table 4 Subclinical atherosclerosis (SCAT)

Coronary artery calcium score (CAC) >10 U Agatston⁴
Carotid plaque (intima-media thickness >1.5 mm) ¹⁴
Computed tomography coronary angiography (CCTA) with a definite plaque ¹⁵
Ankle-brachial index <0.9 ¹⁶
Abdominal aortic aneurysm (AAA)  [17–21]²⁶

⁴ When available, CAC scoring should be the preferred modality

¹⁵ CCTA should not be performed routinely in truly asymptomatic patients

²⁶ Patients suffering from an AAA are at elevated risk of cardiovascular morbidity and mortality, due to common risk factors and comorbidities associated with the aneurysm

Table 5 Clinical atherosclerotic disease (CLAD)

Acute coronary syndrome:
Acute myocardial infarction or unstable angina
Stable angina or previous acute myocardial infarction
Atherothrombotic stroke or transient ischemic attack
Coronary, carotid, or peripheral revascularization
Peripheral vascular insufficiency or limb amputation
Severe atherosclerotic disease (stenosis >50%) in any vascular territory
cardiac infarction, acute coronary syndrome without myocardial infarction, coronary revascularization, or CHD death. The probability of developing CAC by age 32–56 was estimated using clinical risk factors measured 7 years apart between ages 18 and 38. Participants were followed up for 12.5 years, with 57 incident CHD events and 108 incident CVD events observed. After adjusting for risk factors and treatments, those with any CAC had a fivefold increase in CHD events (hazard ratio [HR] 5.0, 95% CI 2.8–8.7) and a threefold increase in CVD events (HR 3.0, 95% CI 1.9–4.7). Within CAC strata of 1–19, 20–99, and >100, the HRs for CHD were 2.6 (95% CI 1.0–5.7), 5.8 (95% CI 2.6–12.1), and 9.8 (95% CI 4.5–20.5), respectively. A CAC score ≥100 was associated with an incidence of 22.4 deaths per 100 participants in 12.5 years (HR 3.7, 95% CI 1.5–10.0). The presence of CAC among individuals aged 32–46 was associated with increased risk of fatal and nonfatal CHD during 12.5 years of follow-up. Thus, screening for CAC might be considered in individuals with risk factors in early adulthood to inform discussions about primary prevention.

- The MESA study was a prospective population-based cohort that investigated the prevalence and progression of subclinical cardiovascular disease in persons without cardiovascular disease at baseline, including 6814 men and women aged 45–84 years and 9.8% with diabetes, to assess the predictive accuracy and improvement in reclassification gained by the addition of CAC score (among others) over the atherosclerotic cardiovascular risk estimator (ASCVD). The authors concluded that CAC score had a modestly improved discriminative ability over ASCVD. The Harrell’s C statistic difference was significant (0.74 vs. 0.76, p = 0.04), and CAC score addition was the only marker that improved ASCVD risk score [30].

3. In patients with diabetes, a CAC score >10 is an indicator of increased mortality and future cardiovascular events. It is recommended that patients with diabetes with a CAC score >10 should be considered as HIGH RISK. [I, A]

Summary of evidence
- In a meta-analysis of eight studies including 6521 patients with diabetes [31], with a mean follow up of 5.18 years, the relative risk of the composite outcome of all-cause mortality and/or cardiovascular events with CAC > 10 vs. CAC < 10 was 5.47 (95% CI 2.59–11.53, p < 0.001) [31]. Nevertheless, it should be noted that significant heterogeneity was detected across studies (I² = 82.4%, p < 0.001). CAC > 10 had a sensitivity of 94% and a specificity of 34% for the composite outcome. A higher CAC score entailed lower sensitivity and higher specificity. For example, when comparing CAC < 10 vs. CAC > 1000, sensitivity dropped to 90%, while specificity increased to 74%. For an individual with diabetes and CAC < 10, post-test probability for the composite outcome was 1.8%, representing a 6.8-fold decrease in the pretest probability of a CVD outcome. The study concluded that a CAC < 10 is helpful to detect lower-risk individuals in this population.
- The Diabetes Heart Study monitored cardiovascular mortality in 1051 patients with diabetes followed for 7.4 years. A positive association was observed between CAC and mortality in the model adjusted for age, sex, race, smoking, and LDL-C. Using the group score of 0–9 for CAC as a reference, the study found the following relative risks according to CAC severity: CAC 10–99: 1.40 (95% CI 0.57–3.74, p = 0.47); CAC 100–299: 2.87 (95% CI 1.17–7.77, p = 0.02); CAC 300–999: 3.04 (95% CI 1.32–7.90, p = 0.008); and CAC ≥ 1000: 6.71 (95% CI 3.09–16.87, p = 0.0001) [32]. Later in 2013, the same authors published an analysis of CAC score compared to traditional risk factors to predict cardiovascular mortality. CAC improved the AUC-ROC from 0.70 (95% CI 0.67–0.73) to 0.75 (95% CI 0.72–0.78). The net reclassification index (NRI) in the moderate-risk group (7–20% in 10 years) was 0.34, which means that 34% of individuals were reclassified into different risk categories [33].

4. CAC score outperforms carotid-artery intima-media thickness (CIMT) and ankle-brachial index (ABI) to discriminate and reclassify cardiovascular risk, at least in nondiabetic subjects. [IIa, B]

Summary of evidence
- The MESA study compared the performance of distinct stratification methods in an intermediate-risk population with no previous cardiovascular event (estimated Framingham risk score between 5 and 20%) [33]. In that study, calcium score (AUC for CAC plus Framingham risk score: 0.784) presented better risk discrimination compared to CIMT (AUC for CIMT plus Framingham risk score: 0.652) and ABI (AUC for ABI plus Framingham risk score: 0.650), as well as better reclassification ability (NRI for calcium score: 0.659; NRI for CIMT: 0.102; NRI for ABI: 0.036) [33]. Although patients with diabetes mellitus were not part of the study, calcium score was shown to be clearly superior to CIMT and ABI to predict risk of coronary events.
5. Carotid plaque can predict major adverse cardiovascular events (MACE) and reclassify risk. Adding plaque information with abnormal wall thickness (CIMT > 1.5 mm) may be useful to reclassify intermediate risk into high risk. [IIb, B]

Summary of evidence
- The Atherosclerosis Risk in Communities (ARIC) study followed 13,145 individuals without previous CVD (57% women, age: 54.0 ± 8.5 years, 10% with diabetes) for a mean of 15.1 years, during which 1812 CHD events occurred [34]. CIMT (categorized as <25th percentile, 25th–75th percentile or >75th percentile for sex) or plaque presence, defined in the presence of at least 2 of 3 criteria—abnormal wall thickness (CIMT > 1.5 mm), abnormal shape (protrusion into the lumen, loss of alignment with adjacent arterial wall boundary), and abnormal wall texture (brighter echoes than adjacent boundaries)—were superior for risk discrimination and reclassification in comparison with risk factors alone. According to the authors, when plaque information (abnormal wall thickness) and CIMT were considered in addition to risk factors, 8.6, 37.5, 38.3, and 21.5% of the overall sample were reclassified in the <5, 5–10, 10–20, and >20% 10-year estimated risk groups respectively. Adding plaque and CIMT reclassified 17.4, 32.8, 36.6, and 25.2% of the men and 5.1, 40.2, 38.4, and 24.9% of the women in the same risk groups.
- The prospective cohort Biolmage Study enrolled 5808 asymptomatic adults without previous cardiovascular events to evaluate the role of vascular imaging in cardiovascular risk prediction [35]. All patients were evaluated for carotid plaque burden score based on a novel 3-dimensional carotid ultrasound and CAC score at baseline, and followed up for a median of 2.7 years. The main study outcome was the presence of MACE defined as cardiovascular death, myocardial infarction, and ischemic stroke. The authors analyzed the carotid plaque burden (cPB) through the sum of the areas of carotid plaques as seen along both carotid arteries and their ramifications. cPB was analyzed in tertiles. After adjustments for risk factors, and compared with individuals without any cPB, hazard ratios (HR) for MACE at tertiles 1, 2, and 3 were 0.78 (95% CI 0.31–1.91), 1.45 (95% CI 0.67–3.14), and 2.36 (95% CI 1.13–4.92) respectively. The net reclassification index (NRI) significantly improved in 23%. Thus, detection of subclinical carotid atherosclerosis improves risk prediction and reclassification compared with traditional risk factors [35].

Module 3: Screening of silent myocardial ischemia
6. A resting electrocardiogram (ECG) should be considered at least annually in asymptomatic patients with diabetes at INTERMEDIATE, HIGH, and VERY HIGH RISK. [IIa, B]

Summary of evidence
- On the basis of expert-opinion evidence, an annual resting ECG is recommended for diabetic patients at high and very high cardiovascular risk, given its low cost, high safety, and prognostic value of ECG abnormalities, which must lead to further exploration.
- In the Epidemiology of Diabetes Interventions and Complications (EDIC) Study [36], patients with type 1 diabetes had a mean follow-up of 19 years and underwent at least one ECG annually. The presence of any major ECG abnormalities was associated with a more than twofold increased risk of CVD events (hazard ratio [HR] 2.10 [95% CI 1.26–3.48] vs. no abnormality/normal ECG, and 2.19 [95% CI 1.46–3.29] vs. no major abnormality).
- In the United Kingdom Prospective Diabetes Study (UKPDS), one in every six newly diagnosed patients with diabetes had ECG evidence of silent myocardial infarction [37].
- The MiSAD study [38] included 925 asymptomatic intermediate to high-risk patients with type 2 diabetes mellitus who underwent an ECG stress test, which, if positive, led to stress myocardial perfusion imaging (MPI). The prevalence of coronary artery disease (CAD) was 12.5% (abnormal exercise test). Of individuals with CAD, 6.4% had abnormal perfusion at MPI. Multivariate analysis showed that, in the overall population, the associated independent risk factors were age, total cholesterol, proteinuria, and, importantly, ST-T abnormalities on resting ECG, which had the highest odds ratio (9.27, CI 4.44–19.38) and was the only risk factor identified in both women and men. Abnormal MPI predicted cardiac events at 5 years (HR 5.5, 95% CI 2.4–12.3, p < 0.001). The relevance of ST-T abnormalities on resting ECG as a predictor of silent CAD highlights the importance of performing periodic resting ECGs in patients with type 2 diabetes.

7. Universal screening for coronary artery disease with stress induction of myocardial ischemia does not improve outcomes and is NOT RECOMMENDED in truly asymptomatic diabetic patients when in the absence of resting ECG abnormalities, even in the presence of a high-risk condition for cardiovascular events. [III, A]

Summary of evidence
- The detection of ischemia in asymptomatic diabetics (DIAD) multicenter randomized trial evaluated
whether detection of silent myocardial ischemia in asymptomatic patients with diabetes could reduce cardiovascular events. The participants were randomized to undergo routine screening for detection of silent ischemia using adenosine stress myocardial perfusion single-photon emission computed tomography (SPECT) or no screening. A total of 1123 asymptomatic diabetic patients were randomized. After a mean follow-up of 4.8 years, a non-significant reduction in the overall cardiac event rate was detected in the screened vs. unscreened group, with HR of 0.88 (95% CI 0.44–1.88) [39].

- A prospective, multicenter randomized trial—do you need to assess myocardial ischemia in type-2 diabetes (DYNAMIT) study [40]—evaluated screening for silent myocardial ischemia using a bicycle exercise test or dipyridamole stress SPECT in 631 asymptomatic diabetic patients with no evidence of coronary artery disease. The study was discontinued prematurely because of difficulties in recruitment and a lower-than-expected event rate. There were no significant differences between the screening and usual-care groups for the main outcome (HR 1.00, 95% CI 0.59–1.71). A meta-analysis of the DYNAMIT and DIAD trials [39] produced similar results, with narrower confidence intervals for each endpoint.

- The BARDOT trial [41] was a prospective multicenter study evaluating the prevalence, progression, treatment, and outcome of silent coronary artery disease (CAD) in 400 asymptomatic patients with diabetes at high coronary risk, without history or symptoms of CAD. Patients underwent myocardial perfusion SPECT (MPS) at baseline and after 2 years [41]. Patients with normal MPS received usual care, while those with abnormal MPS received medical or combined invasive and medical management. An abnormal MPS was found in 22% of patients. Normal-MPS patients had a low rate of first manifestations of CAD compared with patients with abnormal MPS at baseline. Patients with normal MPS had 2-year rates of MACE, cardiac death, and of new ischemia or new scar of 2.9, 0.7, and 3.2% respectively. Patients with abnormal MPS had a sevenfold higher rate of progression to "overt CAD," independent of therapy [41]. However, although the BARDOT trial results suggested screening and treating high-risk patients on the basis of MPS, it is important to note that only about 20% of patients with an abnormal MPS would be advised to receive anti-ischemic therapy. The findings of the Bardot study are preliminary and still require confirmation. A combined medical and invasive strategy may reduce scintigraphic but not symptomatic CAD progression compared with medical therapy alone. Thus, universal screening cannot be currently advised in high-risk patients until more robust data are available.

8. Consider investigation for myocardial ischemia in asymptomatic patients with diabetes when resting ECG abnormalities are present and in patients who exhibit typical or atypical cardiac symptoms (unexplained dyspnea, atypical chest pain or discomfort), evidence of associated vascular disease (carotid bruits, transient ischemic attack, stroke, peripheral arterial disease) and a very high CAC score (>400), when available. [IIa, B]

Summary of evidence

- In a sub-study of the 30-year UKPDS [37], data from 5102 diabetic patients were analyzed through Cox proportional hazards regression to examine outcomes by silent myocardial ischemia (SMI) status. Of 1967 diabetic patients with complete baseline data, 326 (16.6%) had ECG evidence of SMI at enrollment. Around one in six UKPDS patients with newly-diagnosed T2D had evidence of SMI, which was independently associated with an increased risk of fatal MI and all-cause mortality.

- Raggi et al. [26] conducted a 5-year follow-up of 10,377 asymptomatic individuals (903 with diabetes) with a baseline CAC score available. The authors used Cox proportional hazard models, with and without adjustment for other risk factors, to predict all-cause mortality as the primary endpoint. All-cause mortality was increased in asymptomatic patients with diabetes in proportion to the screening CAC. In a risk-adjusted model, there was significant interaction of CAC with diabetes (p < 0.00001), indicating that for every increase in CAC, there was a greater increase in mortality for diabetic compared to nondiabetic subjects. The mortality of diabetic patients with CAC > 400 in the study was around 10% in 4–5 years, greater than that of nondiabetics.

9. Exercise ECG should be considered as the initial test for investigation of ischemia in most symptomatic patients. Exceptions are when resting ECG abnormalities preclude interpretation of exercise stress testing and in patients who are unable to exercise. In those cases, pharmacological stress echocardiography, myocardial perfusion imaging (MPI), coronary computed tomography angiography (CCTA), and stress perfusion cardiac magnetic resonance imaging are reasonable options. [IIa, C]

Summary of evidence

- The treadmill stress test is widely used for CAD detection in the general population because it is easily performed, has relatively good predictive value, and is inexpensive. In diabetic patients, the negative predictive value of the stress ECG is 87%, with 75%
specificity. Lyerly et al. [42] studied 2854 men with documented diabetes mellitus (mean age 49.5 years) who completed a maximal treadmill exercise test with a mean follow-up of 16 years. Those with normal ECG presented the highest CHD-free survival. Those with abnormal ECG and those who were unable to perform maximal exercise had lower CHD-free survival rates. Stress SPECT with thallium or MIBI provides a wide range of information, including ischemia location and extension and left ventricular function, helping physicians appreciate the severity of CAD. This modality can be coupled with pharmacologic agents (dipyridamole, adenosine) for stress induction. In individuals with diabetes, SPECT has higher sensitivity (80–90%) and specificity (75–90%) than the ECG stress test [43]. Another alternative for SMI screening is stress echocardiography using exercise or drugs such as dobutamine. Stress echocardiography detects wall motion abnormalities during stress and provides information on ischemia intensity and left ventricular function. Sensitivity and specificity are 81 and 85% respectively in asymptomatic diabetic patients [44]. CMRI perfusion imaging, with sensitivity of 86.5% and specificity of 83.4% to detect angiographically significant coronary stenosis (>50% left main coronary artery or >70% branch disease), is an alternative for patients who cannot exercise [45].

10. Coronary computed tomography angiography (CCTA) should NOT be used routinely in ASYMPTOMATIC patients with diabetes, since it does not seem to reduce cardiovascular event risk when used for risk stratification of this population. [III, B]

Summary of evidence

- The FACTOR 64 study [46] evaluated whether CCTA was beneficial to reduce clinical outcomes in asymptomatic patients with type 1 or 2 diabetes. Patients with diabetes were included if disease duration was at least 5 years. The patients were randomly assigned to CCTA or optimal diabetes care, and the result of CCTA was used for clinical decision-making. Non-screening patients received standard-of-care treatment for existing risk factors, and physicians were encouraged to reach therapeutic goals in accordance with current guidelines (glycated hemoglobin <7.0%, LDL-c < 100 mg/dL, systolic blood pressure < 130 mmHg). Patients in the screening CCTA arm with normal coronary arteries remained on standard-of-care therapy. Patients who exhibited mild or severe proximal lesions or distal lesions or a CAC score >10 were advised to pursue more aggressive treatment targets (LDL-c < 70 mg/dL, HDL-c > 50 mg/dL, triglycerides <150 mg/dL, glycated hemoglobin <6.0%, and systolic blood pressure <120 mmHg). Patients with severe stenosis underwent invasive coronary angiography, and the decision regarding revascularization was based on the judgment of the assistant physician. Patients with moderate lesions underwent evaluation of myocardial ischemia. Nine hundred patients were randomized, 452 to CCTA, with a mean follow-up of 4 years. Mean duration of diabetes in the group not undergoing CCTA was 13.5 years, vs. 12.3 years in the CCTA arm. The primary endpoint rate (total mortality, nonfatal MI, or unstable angina) was similar, with 28 events (6.2%) in the CCTA group vs. 34 events (7.6%) in the control group (HR 0.80, 95% CI 0.49–1.32, $p = 0.38$). No differences were observed for the secondary endpoint (major ischemic cardiovascular events). In fact, the observed event rate was lower than expected for the sample size, which may explain the negative results. Patients with diabetes in whom risk factors were well controlled did not benefit from CCTA screening as a preventive measure to reduce cardiovascular event risk. Thus, CCTA cannot be recommended for screening of asymptomatic patients with diabetes at this time.

11. In patients at LOW or INTERMEDIATE risk categories, with atypical symptoms, coronary computed tomography angiography (CCTA) may be considered to rule out myocardial ischemia, as it has a good negative predictive value. [IIb, B]

Summary of evidence

- Hadamitzky et al. [47] evaluated the role of CCTA for prediction of cardiovascular events in 140 subjects with diabetes and 1782 without diabetes followed for a mean of 33 months. Participants presented with atypical symptoms of CHD or other risk factors. Those with diabetes and a high plaque burden, as characterized by high number of coronary segments with atherosclerotic plaque (calcified or not), had a much higher event rate than those without diabetes (1.8% vs. 0.5% per year). Plaque burden was the best marker of coronary events, even when adjusted for calcium score.

Module 4: Management of hyperglycemia

Targets

12. In non-pregnant adult patients with type 1 or 2 diabetes mellitus, and in the absence of severe cognitive impairment or reduced life expectancy, the recommended target for glycemic control is a HbA1c below 7.0%. [I, A]

Summary of evidence

- The diabetes control and complications trial (DCCT) [48] and the United Kingdom Prospective Diabetes
Study (UKPDS) [49] demonstrated that achieving an HbA1c below 7% reduces microvascular complications in type 1 and type 2 diabetes. In subjects with type 1 diabetes, implementing intensive glycemic control targeting an HbA1c below 7% in the first 6 years of diabetes can promote a 57% reduction in nonfatal myocardial infarction, stroke, and death from cardiovascular disease on long-term follow-up (9 years), as seen in the DCCT/EDIC study [50, 51]. Similarly, in type 2 diabetes, intensive glycemic control decreases cardiovascular outcomes in the long term (after 10 years) when implemented in recently diagnosed patients [52].

• Lower HbA1c targets were evaluated in three randomized clinical trials: action to control cardiovascular risk in diabetes (ACCORD) [53], action in diabetes and vascular disease: preterax and diamicron modified release controlled evaluation (ADVANCE) [54], and the veterans affairs diabetes trial (VADT) [55]). These trials did not detect reduction in cardiovascular outcomes when intensive control (HbA1c < 6.5%) was implemented. The ADVANCE study (n = 11,140), ACCORD (n = 10,251), and VADT (n = 1791) evaluated patients with type 2 diabetes and previous cardiovascular disease or risk factors and diabetes (mean duration 8–11.5 years), assessing incidence of cardiovascular disease after intensive vs. conventional treatment. The final mean HbA1c was 6.5 vs. 7.3% (ADVANCE), 6.4 vs. 7.5% (ACCORD), and 6.9 vs. 8.4% (VADT). In the ACCORD trial, but not in the other studies, a 22% increase in all-cause mortality followed intensive treatment.

13. Less stringent HbA1c targets (below 8.0%) are reasonable in patients with known history of severe and frequent hypoglycemic events, long-standing disease, short life expectancy, major comorbidities, and established vascular complications, as well as in less motivated, non-adherent patients and in those with diminished self-care capacity, limited resources, and a limited support system. [IIa, B]

Summary of evidence

• Tight glucose control may be harmful in many patients, particularly the elderly and those with other illnesses, especially cardiovascular diseases [56]. Intensive glycemic control does not lead to improved microvascular outcomes for at least 8 years. Data from randomized controlled trials suggest that intensive glycemic control immediately increases the risk of severe hypoglycemia 1.5- to 3-fold [57].

• Observational data from emergency admissions showed a consistent increase in severe hypoglycemia over one decade, especially in type 2 diabetes patients with lower HbA1c, more comedication, and more concomitant diseases [58, 59]. Hypoglycemia in these patients has been associated with increased mortality, higher risk of dementia, falls, fall-related fractures, cardiovascular events, and poor quality of life [60]. Mechanisms by which acute hypoglycemia may trigger ischemia, arrhythmia, and cardiovascular events include increases in epinephrine and norepinephrine levels, which may induce increased cardiac rate and/or contractility, thus heightening myocardial oxygen consumption, while also precipitating vasoconstriction and platelet aggregation. Moreover, acute hypoglycemia in the presence of hypokalemia prolongs cardiac repolarization and increases the QT interval, favoring a proarrhythmic state [60].

• In patients with diabetes from a Brazilian multicenter registry followed for 12 months, failure to reach HbA1c targets was associated with poorer event-free survival (all-cause mortality, nonfatal cardiac arrest, myocardial infarction, or stroke) as compared to good metabolic control (p < 0.041). In that study, HbA1c targets of 8.0 and 7.0% were considered in patients without a previous cardiovascular event vs. those with a previous cardiovascular event [61].

• Patients with limited resources and a limited support system, those with lower motivation, non-adherent patients, and those with diminished self-care capacity are not candidates for strict glucose control, as the risk of hypoglycemia tends to be higher [62].

• Considering the high risk of hypoglycemia with strict metabolic control, especially in elderly patients and in those in which this adverse effect may be more harmful, individualized targets should be pursued in patients with a known history of severe and frequent hypoglycemic events, long-standing disease, short life expectancy, major comorbidities, and established vascular complications [63]. Considering these data and the results of observational studies, the harms associated with an HbA1c target lower than 7.5% or higher than 9% are likely to outweigh the benefits in most adults older than 65 years [57, 64].

• Data to guide this type of individualized treatment are derived from weak evidence. However, the high frequency of risk factors for hypoglycemia and its adverse impact, as well as the marginal benefits of tight control in individuals with short life expectancy, suggest a need to reduce overtreatment, particularly among the elderly and the other groups cited above [56, 60, 64].
14. In hospitalized patients with acute myocardial infarction, it is recommended that blood glucose be maintained in the 130–200 mg/dL range by continuous intravenous insulin infusion, followed by good long-term metabolic control. [I, B]

**Summary of evidence**

- The DIGAMI [65] study included 620 patients with diabetes and acute myocardial infarction (AMI) and used the following strategies: IV infusion of insulin and glucose in the first 24 h with a glycemic target of 126–196 mg/dL, subcutaneous administration of insulin four times daily for 3 months, vs. standard insulin therapy as clinically indicated at the time of the study. Treatment with insulin in the acute phase produced better glycemic control during hospitalization, at 3 months and at 1 year, as well as lower mortality rates at 1 and 3.4 years of follow-up.
- In the DIGAMI-2 trial [66], use of insulin during hospitalization and after discharge was compared to insulin therapy only during hospitalization and usual treatment throughout the period. Glycemic control and cardiovascular outcomes were similar in the two groups.
- In the HI-5 study [67], 240 patients with diabetes and glucose ≥140 mg/dL were included at hospital admission for AMI and randomized to strict glycemic control (target glycemia 72–180 mg/dL) with insulin plus intravenous glucose infusion for at least 24 h or conventional therapy. After discharge, the patients were managed by their physician, with a recommendation to maintain HbA1c < 7%. The groups had similar in-hospital mortality rates.

15. In patients undergoing cardiac surgery, it is recommended that blood glucose be maintained in the 120–150 mg/dL range through continuous intravenous insulin infusion during the hospitalization period. [I, A]

**Summary of evidence**

- Hyperglycemia before or after cardiac surgery has been associated with increased risk of complications (death, prolonged mechanical ventilation, renal failure, stroke, and deep sternal infection) [68, 69].
- The observational Portland Diabetes Project study evaluated the relationship between hyperglycemia and adverse outcomes of cardiac surgery in patients with diabetes. In the study, continuous intravenous insulin, adjusted by frequent blood glucose tests was used based on a standardized protocol conducted by nurses. Initial glucose target was 150–200 mg/dL. This was later changed to 125–175 mg/dL and then to 100–150 mg/dL because other studies were identifying the need to normalize blood glucose reduction outcomes. The use of this protocol vs. subcutaneous insulin according to glucose levels (historical control) was associated with reduced rates of infection [70] and death in about 50% of patients [71].
- A randomized controlled trial with surgical intensive coronary unit patients (63% cardiac surgery and 13% diabetes) showed benefit of intensive glycemic control (insulin infusion glycemic target 80–110 mg/dL) vs. usual glycemic control (180–200 mg/dL) in mortality, infection, acute renal failure requiring hemodialysis, blood transfusion, and polyneuropathy in critically ill patients. However, intensive glycemic control was associated with higher rates of hypoglycemia [72].
- Nevertheless, the multicenter NICE SUGAR Study, conducted in medical (63%) and surgical intensive coronary units (37% of patients respectively; 20% with a history of diabetes), showed that intensive glycemic control (target < 108 mg/dL) vs. usual control (140–180 mg/dL) increased mortality and hypoglycemia rates [73]. A meta-analysis including data from the NICE SUGAR study, with separate analysis of clinical and surgical ICUs, showed that tight glucose control did not reduce mortality in the clinical ICU, but may bring benefit to surgical patients when target blood glucose is <150 mg/dL. [74]. A small RCT comparing two glycemic targets (90–120 mg/dL vs. 120–180 mg/dL) in patients with diabetes undergoing coronary artery bypass grafting showed increased risk of hypoglycemia and absence of benefit with more strict blood glucose control [75].

16. A basal plus bolus correction insulin regimen (a strategy using multiple doses of long- and short-acting insulins) is a reasonable option for correcting hyperglycemia in hospitalized, non-critically ill diabetic patients. [IIa, B] The use of sliding-scale insulin in the inpatient hospital setting is discouraged. [III, C]

**Summary of evidence**

- Hyperglycemia in in-hospital patients with diabetes is very common. Retrospective and randomized controlled trials in surgical populations have reported that hyperglycemia of diabetes is associated with increased length of stay, hospital complications, resource utilization, and mortality [76, 77].
- A randomized controlled trial showed that basal-bolus treatment (glargine and glulisine) improved glycemic control and reduced hospital complications (wound infection, pneumonia, acute renal failure, and bacteremia) compared with sliding-scale insulin (glulisine) in general surgery patients with type 2 diabetes [78].
• Some RCTs were performed in type 2 diabetic patients hospitalized for nonsurgical conditions. In this population, basal–bolus treatment (glargine and glulisine or NPH and regular) also improved glycemic control compared with sliding-scale insulin [79, 80].

Outpatient treatment: monotherapy

17. In patients with recently diagnosed type 2 diabetes, metformin plus non-pharmacological therapy including physical activity and targeted nutrition therapy for weight control is recommended as first-line therapy. [I, A]

Summary of evidence
• Metformin has a favorable efficacy and safety profile, with important metabolic effects and cardiovascular benefits. Due to its effect in reducing cardiovascular events and mortality, its efficacy in blood glucose reduction with low incidence of hypoglycemia, low cost, tolerable adverse effects, and no association with weight gain, it is the current first-line agent of choice for treatment of hyperglycemia in type 2 diabetes [81]. Titration or addition of further hypoglycemic drugs should be implemented as soon as possible to avoid inertia in achieving glucose targets.

18. In patients who do not tolerate metformin, any other antidiabetic drug can be recommended as monotherapy, except if contraindicated. [I, C]

Summary of evidence
• The UKPDS analyzed 5102 recently diagnosed type 2 diabetes patients followed up from 1977 to 1997 and found that intensive glycemic control with sulfonylurea or insulin therapy decreases progression of microvascular disease and may also reduce the risk of heart attacks. In obese patients, the UKPDS showed that metformin has similar efficacy to sulfonylureas for glucose control [52, 82, 83].

• UKPDS 34 investigated whether intensive glucose control with metformin has any specific advantage or disadvantage. A subgroup analysis compared 411 recently diagnosed overweight (>120% ideal bodyweight) type 2 diabetes patients treated with diet alone versus 342 patients using metformin, aiming for a fasting plasma glucose <110 mg/dL, and found a relative risk reduction (RRR) of 32% ($p = 0.002$) of any diabetes-related complications, a 42% RRR for any death related to diabetes ($p = 0.017$), and a 36% RRR for all-cause mortality ($p = 0.011$).

Table 6 Renal function adjustments of anti-hyperglycemic drugs

| Drug            | Maximal daily dose | Estimated GFR (mL/min) | 45–60 | 30–45 | <30 |
|-----------------|--------------------|------------------------|-------|-------|-----|
| Insulin         | Variable           | NN                     | NN    | NN    | NN  |
| Pioglitazone    | 45 mg              | NN                     | NN    | NN    | NN  |
| Linagliptin     | 5 mg               | NN                     | NN    | NN    | NN  |
| Sitagliptin (mg)| 100                | 50                     | 50    | 50    | 25  |
| Vildagliptin    | 50 mg bid          | 50 mg                  | 50 mg | 50 mg | 50 mg |
| Saxagliptin (mg)| 5                  | 2.5                    | 2.5   | 2.5   | 2.5 |
| Albiglaptin (mg)| 25                 | 12.5                   | 12.5  | 12.5  | 6.25|
| Metformin       | <2550 mg           | <2000 mg/day           | <1000 mg/day | NN |
| Glimipride      | 8 mg               | 1 mg                   | 1 mg  | 1 mg  | NR  |
| Gliclazide      | 120 mg             | NN                     | NN    | NN    | NR  |
| Glibenclamide   | 20 mg              | NR                     | NR    | NR    | NR  |
| Nateglinide     | 120 mg/meal        | 60 mg/meal             | 60 mg/meal | NR |
| Repaglinide (mg/meal)| 1          | 0.5                    | 0.5   | 0.5   | NR  |
| Acarbose        | 300 mg             | 150 mg                 | 150 mg| 150 mg| NR  |
| Exenatide       | 10 mcg bid         | NN                     | 5 mg  | NR    | NR  |
| Liraglutide     | 1.8 mg             | NN                     | NN    | NN    | NR  |
| Lixisenatide    | 20 mcg             | NN                     | NN    | NR    | NR  |
| Dulaglutide     | 1.5 mg/week        | NN                     | NN    | NR    | NR  |
| Canagliflozin   | 300 mg             | 100 mg                 | 100 mg| NR    | NR  |
| Empagliflozin   | 25 mg              | NN                     | NN    | NR    | NR  |
| Dapagliflozin   | 10 mg              | NN                     | NN    | NR    | NR  |

GFR glomerular filtration rate, NN not necessary, NR not recommended, bid 2 times daily

19. In patients with renal impairment, possible substitutions of anti-hyperglycemic drugs for type 2 diabetes are indicated in Table 6

Outpatient treatment: second agent

20. In an asymptomatic patient with recently diagnosed type 2 diabetes and HbA1c > 8.5%, combined pharmacological treatment for hyperglycemia consisting of metformin plus a second antihyperglycemic agent should be considered as first-line therapy. [IIa, C]

Summary of evidence
• This is an expert opinion-based recommendation, not based on published evidence. The majority of members from the Panel recommends to start combined therapy with metformin above HbA1c > 8.5% to avoid delaying the attainment of optimal glycemic control; all efforts should be made to prevent severe hyperglycemia in treatment-naïve patients with type 2 diabetes.
21. In patients who do not achieve target HbA1c levels on monotherapy, any antidiabetic drug is potentially effective as an add-on option to metformin for glycemic control. There is no evidence of significant differences between classes of antidiabetic agents when used as second therapy added to metformin. [I, A] Pharmacological therapy to lower blood glucose in the patient with type 2 diabetes should be individualized on the basis of efficacy, mechanism of action, presence of comorbidities, risk of hypoglycemia, weight gain, adverse effects, and costs. [I, C]

Summary of evidence

- A meta-analysis [84] including 27 RCTs with 11,198 type 2 diabetes patients showed a similar HbA1c reduction between different classes of antidiabetic agents compared to placebo. The mixed-treatment comparison showed the following reductions in HbA1c: sulfonylureas, 0.79%; glinides, 0.65%; thiazolidinediones, 0.85%; α-glucosidase inhibitors, 0.64%; DPP-4 inhibitors, 0.78%; and GLP-1 analogues, 0.97%. Thiazolidinediones, sulfonylureas, and glinides were associated with mild weight gain, while GLP-1 analogues were associated with a significant decrease in body weight compared with placebo (−1.74 kg). There was no weight change associated with α-glucosidase inhibitors or DPP-4 inhibitors. Sulfonylureas and glinides were associated with an increased risk of hypoglycemia compared with placebo [84]. The choice of second antidiabetic agent should be based on efficacy, age, mechanism of action, risk of hypoglycemia, presence of comorbidities, life expectancy, weight gain or loss, adverse effects, and potential for cardiovascular protection [85].

Outpatient treatment: third agent

22. There is also no difference in HbA1c reduction when different classes of drugs are used as a third option for the treatment of type 2 diabetes. This panel recommends that any antidiabetic drug could be an option as a third agent for glycemic control, provided that the mechanism of action is not similar to that of agents already in use. [I, C]

Summary of evidence

- A meta-analysis of 18 RCTs (n=4535) evaluated the comparative efficacy of GLP-1 agonists, DPP-4 inhibitors, thiazolidinediones, and α-glucosidase inhibitors in reducing HbA1c, body weight, and causing severe hypoglycemia when a third drug was added to a metformin plus sulfonylurea regimen [86]. Despite limitations, as most of the studies were of short duration, with variable quality, and based on indirect comparisons, the meta-analysis showed that all antidiabetic classes were associated with significant reductions in HbA1c levels compared to placebo. The overall average reduction in Hba1c was −0.96% (thiazolidinediones, −1.15%; acarbose, −0.6%; GLP-1 agonists, −1.04%; DPP-4 inhibitors, −0.89%). There was no clear difference in efficacy between drug classes when adding a third agent to treatment of patients with type 2 diabetes who were already receiving metformin and a sulfonylurea [86].

23. Insulin therapy (with or without additional agents) should be considered any time in patients with type 2 diabetes who present persistently high blood glucose levels despite antidiabetic agent combinations, or in patients who are markedly symptomatic. [I, C]

Summary of evidence

- After at least 3 months using metformin plus a second antidiabetic agent, if the glycemic target is not reached, a third drug should be chosen, taking into account the established therapeutic target, age, patient limitations, and the attributes and side effects of each drug. Consider initiating insulin therapy (with or without additional agents) in patients with type 2 diabetes who remain markedly symptomatic (weight loss, ketosis, polyuria, or polydipsia) and/or exhibit elevated blood glucose levels or HbA1c [85].

24. Insulin is a safe option for glycemic control in type 2 diabetic patients treated with one or more antidiabetic agents who do not achieve HbA1c targets or who have typical symptoms of hyperglycemia, even in the presence of high cardiovascular risk. [I, A]

Summary of evidence

- The UKPDS [52] and ORIGIN [87] are randomized controlled trials that used human insulin and the insulin analogue glargine, respectively, in type 2 diabetes and evaluated long-term cardiovascular outcomes. The UKPDS revealed a 15% reduction in myocardial infarction and a 13% reduction in death among people with new-onset type 2 diabetes treated intensively with antidiabetic agents and insulin, as needed to attain an HbA1c of 7.0% vs. usual care. The mean follow up was of 10 years [52].

- In the ORIGIN study [87], participants were randomly assigned to insulin glargine added as an evening injection to their preexisting anti-hyperglycemic regimen or to standard care (treatment according to the investigator’s discretion in alignment with local guidelines). The study included 12,537 people, 88% of whom with type 2 diabetes, of which 59% had a previous cardiovascular event. After a mean follow up of 6.2 years, no differences were found between groups concerning the composite endpoint of nonfatal myocardial infarction, non-
fatal stroke or cardiovascular death. These data indicate that basal insulin treatment (human insulin or insulin analogues) is safe in individuals with type 2 diabetes with or without pre-existing cardiovascular events.

- In the DEVOTE study (Efficacy and Safety of degludec versus glargine in type 2 diabetes), 7637 patients with type 2 DM were randomized to receive either insulin degludec or insulin glargine U100. The primary outcome (nonfatal myocardial infarction, nonfatal stroke and cardiovascular death) occurred in 8.5% of the patients treated with degludec and in 9.3% of the patients treated with glargine (hazard ratio = 0.91; \( p = \text{non-significant} \)). Patients treated with degludec experienced significant lower rates of severe hypoglycemia in comparison to the glargine U100 group (\( p < 0.001 \)) [88].

**Cardiovascular risk**

25. In type 2 diabetic patients at VERY HIGH RISK (presence of clinical atherosclerotic disease, with previous cardiovascular events), the addition of an SGLT-2 inhibitor with demonstrated cardiovascular benefit can be useful to reduce cardiovascular risk, as it reduces the incidence of cardiovascular events and hospitalization due to heart failure in this population. [IIa, A]

**Summary of evidence**

- The EMPA-REG study of empagliflozin, an inhibitor of sodium glucose co-transporter-2 (SGLT2), evaluated 7020 high-risk patients with type 2 diabetes. After 3.1 years, empagliflozin therapy was associated with a 14% reduction in the composite primary outcome of CV mortality, nonfatal AMI, and nonfatal stroke (10.5% vs. 12.1%, \( p = 0.04; \text{NNT} 62 \)), as well as a reduction in all-cause mortality (5.7% vs. 8.3%, \( p < 0.001 \); RRR –32%, NNT 38). There was also a reduction in cardiovascular mortality (3.7% vs. 5.9%, \( p < 0.001 \); RRR –38%, NNT 45) [89]. Interestingly, the HbA1c reduction with empagliflozin was modest (0.5%). The mechanisms by which the drug may have led to this significant result are still being studied.

- The CANVAS Program (Canagliflozin and Cardiovascular and Renal Events in Type 2 Diabetes) included 10,142 patients with type 2 DM, including individuals with established cardiovascular disease (secondary prevention) and patients at high risk for CV events (primary prevention). Patients were then randomized for Canagliflozin (100 mg and 300 mg) or placebo, and were followed for a mean of 188.2 weeks. Canagliflozin therapy was associated with a 14% reduction in the composite primary outcome of CV mortality, nonfatal AMI, and nonfatal stroke (occurring in 26.9 vs. 31.5 participants per 1000 patients-years). However, patients receiving canagliflozin experienced a significant increase in rates of amputation (6.3% vs. 3.4%; \( p < 0.001 \)) and bone fractures (15.4% vs. 11.9%; \( p = 0.02 \)) [90].

- Both EMPA-REG and CANVAS demonstrated a significant reduction in a secondary endpoint composed of hospitalization for heart failure and cardiovascular death.

26. In type 2 diabetic patients with clinical atherosclerotic disease (CLAD) (i.e., VERY HIGH-RISK patients), the addition of a GLP-1 analogue with demonstrated cardiovascular benefit may be useful to reduce cardiovascular risk, as it seems to decrease the incidence of cardiovascular events in this population. [IIa, A]

**Summary of evidence**

- The LEADER study of liraglutide, a GLP-1 analogue, assessed 9340 type 2 diabetes patients with high cardiovascular risk profile. After 3.8 years of follow-up, liraglutide was associated with a 13% reduction in the composite primary outcome of CV mortality, nonfatal AMI, and nonfatal stroke (13% vs. 14.9%, \( p = 0.01 \)). There were reductions in cardiovascular mortality (4.7% vs. 6%, \( p = 0.007 \); RRR –22%) and all-cause mortality (8.2% vs. 9.6%; \( p = 0.02 \); RRR –15%). There was no reduction in the incidence of nonfatal AMI, nonfatal stroke, or hospitalization for heart failure [91].

- The SUSTAIN-6 trial analyzed 3297 patients with longstanding type 2 diabetes (mean disease duration 13.9 years) and established cardiovascular disease, chronic kidney disease, or both, on a standard care regimen, who were randomly assigned to receive once-weekly semaglutide (0.5 or 1.0 mg) or placebo for 104 weeks. At 2-year follow-up, there was a 26% reduction in the composite primary outcome or CV mortality, nonfatal AMI, and nonfatal stroke (6.6% vs. 8.9%, \( p = 0.02 \)) [92]. Cardiovascular death was similar in the two groups (\( p = 0.92 \)). Nonfatal stroke was the main composite primary outcome driver (1.6% vs. 2.7%, \( p = 0.04 \); RRR –39%). Diabetic retinopathy was more frequent in the semaglutide group (3%) than the placebo group (1.8%) (HR 1.76, 95% CI 1.11–2.78, \( p = 0.02 \)). How much of this is due to a greater decrease in HbA1c still needs to be clarified (1% difference between semaglutide 1 mg and placebo).

27. In type 2 diabetic patients, at any level of risk of cardiovascular events, pioglitazone, DPP4 inhibitors, or GLP-1 analogues are safe and reasonable options to achieve glycemic control. [I, A]

**Summary of evidence**

- The use of pioglitazone in patients with long term type 2 diabetes and preexisting CV disease margin-
ally reduced fatal and nonfatal myocardial infarction when compared to placebo (RRR −16%, 95% CI 0.72–0.98, p < 0.03). However, there was a twofold risk of hospitalization for heart failure and an increased risk of bone fractures in women, but no increase in mortality risk [93].

- Recently, several DPP-4 inhibitors and GLP-1 analogues have been evaluated for global CV safety and mortality outcomes in patients with type 2 diabetes at high risk of CV events.
- The TECOS (sitagliptin) study enrolled 14,671 patients with longstanding type 2 diabetes (mean disease duration 11.6 years), preexisting CV disease, and a mean baseline HbA1c of 7.2% [94].
- The SAVOR-TIMI 53 (saxagliptin) study examined 16,492 patients with longstanding type 2 diabetes (mean disease duration 10.3 years), preexisting CV disease or multiple risk factors, and an average baseline HbA1c of 8% [95].
- The EXAMINE (alogliptin) study evaluated 5380 patients with type 2 diabetes (mean disease duration 7.2 years) associated with acute coronary syndrome and average baseline HbA1c of 8% [96].
- The ELIXA (lixisenatide) study examined 6068 patients with type 2 diabetes (mean disease duration 9.4 years) associated with preexisting coronary artery disease with a recent hospital admission due to acute coronary syndrome and mean baseline HbA1c of 7.6% [97].
- Importantly, these studies were designed for noninferiority and demonstrated neutrality regarding global CV safety in patients with type 2 diabetes at high risk of CV events. Saxagliptin was associated with an unexpected increase in hospitalization for heart failure [94–97].

28. In type 2 diabetic patients at any level of cardiovascular risk, the use of sulfonylureas is safe and a reasonable option to achieve glycemic control. However, careful use of sulfonylureas is advocated because of a possible increased risk of hypoglycemia (especially in the elderly), as well as weight gain. [Ia, B]

**Summary of evidence**

- A meta-analysis of 47 RCTs (n = 37,650) evaluated the safety of the most frequently used sulfonylureas, in an attempt to elucidate conflicting data regarding the safety of this class of antidiabetics in terms of mortality and cardiovascular outcomes. The result showed that sulfonylureas were not associated with all-cause mortality (RR 12%, 95% CI 0.96–1.30) or cardiovascular mortality (RR 12%, 95% CI 0.87–1.42). Sulfonylureas were not associated with increased risk of myocardial infarction (RRR −8%, 95% CI 0.76–1.12) or stroke (RR 16%, 95% CI 0.81–1.66) [98].
- The Action in Diabetes and Vascular Disease: Preterax and Diamicron Modified Release Controlled Evaluation (ADVANCE) study and the ADVANCE-ON post-trial study were the largest (and with the highest CV risk population) ever conducted in patients with diabetes on sulfonylurea therapy in which cardiovascular outcomes were determined. The ADVANCE trial randomly assigned 11,140 patients with type 2 diabetes, of whom 32% had pre-existing cardiovascular disease, to undergo either standard glucose control or intensive glucose control, defined as the use of gliclazide MR 60 mg/day to 120 mg/day, plus other drugs, to achieve an HbA1c value of 6.5% or less. After a median of 5 years of follow-up, the mean HbA1c level was lower in the intensive-control group (6.5%) than in the standard-control group (7.3%). Intensive control reduced the incidence of combined major macrovascular and microvascular events (18.1%, vs. 20.0% with standard control; HR 0.90, 95% CI 0.82–0.98, p = 0.01). This was due primarily to a reduction in the incidence of nephropathy (4.1% vs. 5.2%; HR 0.79, 95% CI 0.66–0.93, p = 0.006). Importantly, there was no increase in death from all causes (p = 0.91) nor from cardiovascular causes (p = 0.63). The ADVANCE-ON study invited 8944 surviving participants from the ADVANCE study to a 6-year post-trial study, defining death from any cause and major macrovascular events as primary endpoints. Between-group differences in HbA1c levels during the trial were no longer evident. No differences were observed in risk of death from any cause (p = ) or major macrovascular events between the intensive-control group and the standard-control group (HR 1.00, 95% CI 0.92–1.08 and HR 1.00, 95% CI 0.92–1.08 respectively).

**Module 5: Management of dyslipidemia**

29. In patients with diabetes at VERY HIGH RISK, the recommended lipid target is to reduce LDL-c to a level below 50 mg/dL or non-HDL-c to a level below 80 mg/dL (Table 7). For patients not on statin treatment, at any baseline LDL-c level, an initial reduction in LDL-c or in non-HDL-c of more than 50% from baseline is recommended. [I, A]

**Summary of evidence**

- Two double-blind, controlled, randomized clinical trials have demonstrated that reducing the levels of
LDL-c cholesterol to below (or near) 50 mg/dL is associated with a significant reduction in the incidence of major cardiovascular events. In the FOURIER trial [99], 27,564 patients with atherosclerotic cardiovascular disease and under statin therapy were randomized to placebo or evolocumab. Patients randomized to evolocumab had their LDL-c levels reduced to 30 mg/dL and had a significant reduction in major cardiovascular events (9.8% in the evolocumab group vs 11.3% in the placebo group, hazard ratio 0.85, \( p < 0.001 \)). In the IMPROVE-IT trial [100], 18,144 patients who had been hospitalized for acute coronary syndrome in the preceding 10 days were randomized to simvastatin or simvastatin + ezetimibe. Patients randomized to simvastatin plus ezetimibe had their LDL-c levels reduced to 53.7 mg/dL and experienced a significant reduction in cardiovascular events (32.7% for the simvastatin/ezetimibe group vs 34.7% for the simvastatin group, hazard ratio 0.936, \( p = 0.016 \)).

- Statins have largely been proven to reduce the risk of cardiovascular events in patients with diabetes with a previous history of vascular events. A meta-analysis of 14 trials including 18,686 patients with diabetes concluded that statin treatment reduces the incidence of vascular events proportionately by 20% for each 39 mg/dL reduction in LDL-c in 5 years, with a similar reduction for major coronary events, stroke, and need for revascularization [101].

- In a meta-analysis of individual data from 8 statin RCTs [102] including 38,153 patients allocated into statin therapy, in which lipids and apolipoproteins were determined at baseline and after 1 year of follow-up, a total of 6286 major cardiovascular events were observed in 5387 study participants. Patients with LDL-c below 50 mg/dL were at significantly lower risk than patients with increased levels of LDL-c. The risk category was proportionally lower as the level of LDL-c decreased. Compared with patients whose LDL-c was >175 mg/dL, those who reached an LDL-c of 75–100 mg/dL, 50–75 mg/dL, and <50 mg/dL respectively had progressively lower adjusted HRs of 0.56 (95% CI 0.46–0.67), 0.51 (95% CI 0.42–0.62), and 0.44 (95% CI 0.35–0.55) for major cardiovascular events. Similar associations were observed for non-HDL-c and apolipoprotein B. LDL-c limits may be transferred to non-HDL-c limits by adding 30 mg/dL [103].

- Non-HDL-c is calculated by subtracting HDL-c from total cholesterol. This measure is not affected by triglyceride concentration and is better than calculated LDL-c in patients with increased plasma triglyceride concentrations.

31. **Patients with diabetes in the VERY HIGH-RISK category should initiate statins as soon as possible at the highest tolerable dose (Table 8) to meet cholesterol targets (Table 7).** The lipid profile should be reviewed every 1–3 months. If targets are not met, intensification of treatment is advised, either by switching to a more potent statin, increasing statin dose, adding ezetimibe, and/or improving lifestyle modifications. [I, A]

**Summary of evidence**

- A pre-specified subgroup analysis of the treat to new targets (TNT) study [104], which included 1501 patients with diabetes and coronary artery disease, compared the impact of atorvastatin 80 mg vs. 10 mg on cardiovascular outcomes during 4.9 years. The study showed a significant reduction in any cardiovascular event and stroke in the 80 mg arm. The lower LDL-c attained with the highest dose showed additional benefit.

| Level of risk | Off statin treatment | On statin treatment |
|---------------|----------------------|---------------------|
|               | % Reduction          | LDL-c (mg/dL)       | Non-HDL-c (mg/dL) |
| LOW           | 30–50                | <100                | <130                |
| INTERMEDIATE  | 30–50                | <100                | <130                |
| HIGH          | >50                  | <70                 | <100                |
| VERY HIGH     | >50                  | <50                 | <80                 |

**Table 7** Cholesterol targets in patients with diabetes
PCSK9 inhibitors, however, must be carefully evaluated despite high intensity statin use. The decision to use HIGH-RISK patients who do not meet LDL-c targets 32. The use of PCSK9 inhibitors may be considered in VERY HIGH-RISK patients who do not meet LDL-c targets despite high intensity statin use. The decision to use PCSK9 inhibitors, however, must be carefully evaluated through cost-benefit analysis. [Ila, B] • Monoclonal antibody inhibitors of proprotein convertase subtilisin–kexin type 9 serine protease (PCSK9), a protein that regulates the recycling of LDL receptors, have recently been approved by the FDA, EMEA, and ANVISA for primary prevention in patients with hetero- and homozygous familial hypercholesterolemia or as secondary prevention in patients with CLAD who require additional LDL-c–lowering therapy. This class of drugs meets a large unmet need for more aggressive lipid-lowering therapy beyond statins in an attempt to further reduce residual risk in many persons with clinical CLAD and diabetes. When added to maximal statin therapy, these once- or twice-monthly injectable agents reduce LDL-c by approximately 60%, and have favorable effects on other lipids [106–112]. In post hoc cardiovascular safety analyses of alirocumab and evolocumab added to statins with or without other lipid-lowering therapies, mean LDL-c levels of 48 mg/dL were associated with statistically significant relative risk reductions of 48–53% in major CLAD events [107, 108]. Furthermore, a subgroup analysis of patients with diabetes taking alirocumab demonstrated that a 59% LDL-c reduction was associated with a CLAD event relative risk reduction trend of 42% [113]. • The FOURIER study [99] was a randomized, double-blind, placebo controlled trial which evaluated whether the PCSK9 inhibitor evolocumab associated with a statin could reduce cardiovascular risk vs. statin therapy alone in patients with clinically evident atherosclerotic cardiovascular disease and LDL-c levels of 70 mg/dL. After 48 weeks, evolocumab reduced LDL-c from a baseline of baseline of 92–30 mg/dL and met its primary composite endpoint, reducing cardiovascular death, nonfatal MI or nonfatal stroke (95% CI 0.74–0.95, p < 0.005). Moderate treatment promoted a 30% decrease in cardiovascular events compared to placebo. Intensive treatment promoted a 20% reduction in cardiovascular events beyond moderate treatment. Thus, the overall reduction in events with intensive treatment compared to moderate treatment was 50%. • Treatment goals, even with the highest tolerated statin dose, may not be reached by patients with dyslipidemia, particularly those with established CVD, DM, or asymptomatic high-risk individuals. In such cases, combination treatment may be needed. However, the only combination with evidence of clinical benefit (one large RCT) is that of a statin and with ezetimibe [100]. Based on the relatively limited evidence, the ESC/EAS 2016 panel recommends restricted use of this combination in patients at high or very high risk of CVD [103].

| Statin            | Mean expected LDL-c reduction (%) |
|-------------------|----------------------------------|
|                   | <30 (mg) | 30–50 (mg) | ≥50                  |
| Simvastatin       | 10       | 20–40      | 40 mg + ezetimibe    |
| Pravastatin       | 10–20    | 40–80      | –                   |
| Fluvastatin       | 20–40    | 80         | –                   |
| Atorvastatin      | –        | 10–20      | 40–80 mg            |
| Rosuvastatin      | –        | 5–10       | 20–40 mg            |
| Pitavastatin      | 1        | 2–4        | –                   |
| Lovastatin        | 20       | 40         | –                   |

Table 8 Mean expected % of LDL-c reduction with statin use

- A meta-analysis of five randomized trials [105] (39,612 subjects with prior vascular disease, 5639 [14%] with diabetes) compared intensive vs. moderate statin treatments. Mean follow-up was of 5.1 years. Intensive treatment was defined as a reduction in LDL-c of 20 mg/dL beyond the result obtained by moderate treatment with the use of higher-potency statins. The results showed a 15% further reduction in major vascular events (95% CI 11–18, p < 0.0001), 13% in coronary death (95% CI 7–19, p < 0.0001), 19% in coronary revascularization (95% CI 15–24), p < 0.0001) and 16% in stroke (95% CI 0.74–0.95, p = 0.005). Moderate treatment promoted a 30% decrease in cardiovascular events compared to placebo. Intensive treatment promoted a 20% reduction in cardiovascular events beyond moderate treatment. Thus, the overall reduction in events with intensive treatment compared to moderate treatment was 50%.

32. The use of PCSK9 inhibitors may be considered in VERY HIGH-RISK patients who do not meet LDL-c targets despite high intensity statin use. The decision to use PCSK9 inhibitors, however, must be carefully evaluated through cost-benefit analysis. [Ila, B]
33. In patients with diabetes at VERY HIGH RISK (Tables 2, 5) with a recent acute coronary syndrome, lipid profile should be determined in the first 12–24 h of hospitalization to define baseline levels. Subsequently, statin treatment should be started at the highest tolerable doses, as soon as possible, for 3 months, independently of lipid levels. At that time, lipid profile should be reassessed to check for target achievement. [I, B]

Summary of evidence

- The double-blind, randomized IMPROVE IT trial [100] studied 18,144 patients who had been hospitalized for acute coronary syndrome within the preceding 10 days and had LDL-c levels of 50–100 mg/dL while on lipid-lowering therapy or 50–125 mg/dL if they were not receiving lipid-lowering therapy. The combination of simvastatin (40 mg) and ezetimibe (10 mg) (simvastatin–ezetimibe) was compared with simvastatin (40 mg) and placebo. The primary outcome was the composite of cardiovascular death, nonfatal myocardial infarction and unstable angina requiring re-hospitalization, coronary revascularization, or nonfatal stroke. The median follow-up was 6 years. The median time-weighted mean LDL-c level during the study was 53.7 mg/dL in the simvastatin–ezetimibe group, as compared with 69.5 mg/dL in the simvastatin alone group (p < 0.001). The event rate for the primary endpoint at 7 years was 32.7% in the simvastatin–ezetimibe group, as compared with 34.7% in the simvastatin-alone group, with an absolute risk difference of 2.0% (HR 0.936, 95% CI 0.89–0.99, p = 0.016). Thus, vigorous reduction of LDL-c in the early phases of acute coronary syndrome resulted in improved cardiovascular outcomes and should be recommended. In addition, subgroup analysis revealed the greatest benefit in patients with diabetes, with a 15% reduction in the primary endpoint and a NNT = 18 [100].

34. In patients with diabetes at HIGH RISK (Tables 2, 3, 4), LDL-c should be maintained below 70 mg/dL and/or non-HDL-c below 100 mg/dL. [I, A]
therapy on carotid intima media thickening (CIMT) progression. Statin therapy was found to slow the progression of carotid atherosclerosis, indicating benefits at the subclinical stage of the disease process [117].

36. In patients with diabetes at HIGH RISK, with either stratifying factors (Table 3) or confirmed subclinical atherosclerosis (Table 4), it is highly recommended to start statin therapy (Table 8) to meet targets (Table 7). [I, A]

37. If, after 3 months, LDL-c or non-HDL-c is not at the defined target, intensification of therapy should be considered. [IIa, B]

Summary of evidence
- In the CTT meta-analysis [105] moderate treatment promoted a 30% decrease in cardiovascular events compared to placebo. Intensive treatment promoted a 20% reduction in cardiovascular events beyond moderate treatment. Thus, there was an overall 50% reduction in events with intensive treatment compared to moderate treatment. Despite the indirect evidence provided by subgroup analysis of diabetic patients in the meta-analysis, the absence of heterogeneity makes these results applicable to patients with DM in primary prevention.

38. In patients with diabetes at LOW-INTERMEDIATE RISK, LDL-c levels should be lowered and maintained below 100 mg/dL and non-HDL-c levels should be lowered and maintained below 130 mg/dL (Table 7). [I, B]

Summary of evidence
- In a meta-analysis of 14 trials including 18,686 individuals with diabetes, statin therapy reduced all-cause mortality and vascular mortality, and the reduction in vascular events was proportional to the LDL-c reduction. The proportional effects of statins in diabetic patients were similar irrespective of prior history of vascular disease or other baseline clinical conditions [101].

39. Statins are initially optional for LOW RISK patients, but should be considered in INTERMEDIATE RISK patients (Table 9), if LDL-c and non-HDL-c are above the targets (Table 7). Lipid profile should be re-checked periodically to ensure that LDL-c level is below 100 mg/dL. Intensification of treatment is needed if targets are not met. [IIa, C]

Summary of evidence
- The TRIALIST meta-analysis [118] compared the effects of lowering cholesterol with statins on the incidence of cardiovascular events in a low-risk population. The meta-analysis included 22 statin vs. control trials (n = 134,537) with mean follow-up duration of 4.8 years, and five more vs. less statin trials (n = 39,612) with 5.1 years of follow-up. Participants were separated into five categories of baseline 5-year major vascular event risk on control therapy (<5, ≥5 to <10, ≥10 to <20, ≥20 to <30, ≥30%), with estimation of the rate ratio (RR) per 1.0-mmol/L LDL-c reduction in each category. Reduction of LDL cholesterol with a statin reduced the risk of major vascular events (RR 0.79, 95% CI 0.77–0.81, per 1.0 mmol/L reduction), irrespective of age, sex, baseline LDL-c, or previous vascular disease, and of vascular and all-cause mortality. The proportional reduction in major vascular events was at least as great in the two lowest risk categories as in the higher risk categories. This reflected significant reductions in major coronary events in the two lowest risk categories. In individuals with 5-year risk of major vascular events <10%, each 1-mmol/L reduction in LDL-c produced an absolute reduction in major vascular events of about 11 per 1000 over 5 years. These data indicate that low-risk populations also benefit from lowering cholesterol with statins.

40. It is recommended that patients with diabetes and LDL-c > 190 mg/dL be investigated for familiar hypercholesterolemia (FH). [I, C]

Summary of evidence
- The diagnosis of FH in patients with diabetes should be always considered and further investigated when an LDL-c level > 190 mg/dL is found [106]. LDL-c > 250 mg/dL in a patient aged 30 or older, LDL-c > 220 mg/dL in patients aged 20–29, and LDL-c > 190 mg/dL in patients under age 20 yields approximately 80% probability of FH in the setting of general population screening [107].

41. It is recommended that patients with diabetes and chronic kidney failure who are on dialysis, without CLAD (Table 5), do NOT initiate use of statins, since there is no evidence of benefit in this population and, in fact, the risk of stroke may increase. [III, A] However, in patients with chronic renal failure who were already on statin therapy before initiation of dialysis, withdrawal of statins is not recommended. [III, A]

Summary of evidence
- In the 4D (die deutsche diabetes dialyze) study [119], 1255 patients with type 2 diabetes on hemodialysis were evaluated. They were randomized to atorvastatin 20 mg or placebo and followed up for 4 years. The primary endpoint was a composite of death from cardiac causes, nonfatal myocardial infarction, and stroke. A 42% reduction in LDL-c was observed in
patients on atorvastatin, with no reduction in the primary outcome. The risk of stroke was also increased in this group.

- The study to evaluate the use of rosuvastatin in subjects on regular hemodialysis (AURORA) study [120] included 2776 hemodialysis patients (aged 50–80, 27.9% with diabetes) treated with rosuvastatin 10 mg/day or placebo during a mean of 3.8 years. The primary outcome was a composite of nonfatal myocardial infarction, nonfatal stroke, and cardiovascular death. There was a 43% reduction in LDL-c in the intervention group, but no differences in the primary outcome were observed between groups.

- Regarding patients with chronic renal disease but not on hemodialysis, the Pravastatin Pooling Project database made a combined analysis of results of three randomized trials of pravastatin 40 mg vs. placebo [121], including 19,700 patients with chronic renal insufficiency (estimated GFR 60–30 mL/min/1.73 m²). Significant benefit of treatment was detected in reducing the primary endpoint of myocardial infarction, coronary death, or percutaneous revascularization and total mortality in this group of patients.

- The SHARP trial aimed to assess the efficacy and safety of the combination of simvastatin plus ezetimibe in people with moderate-to-severe kidney disease. This randomized, double-blind trial included 9270 patients with chronic kidney disease (3023 on dialysis and 6247 not on dialysis) with no known history of myocardial infarction or coronary revascularization. Patients were randomly assigned to simvastatin 20 mg plus ezetimibe 10 mg daily versus matching placebo. The key pre-specified outcome was first major atherosclerotic event (nonfatal myocardial infarction or coronary death, non-hemorrhagic stroke, or any arterial revascularization procedure). All analyses were by intention to treat. A total of 4650 patients were assigned to receive simvastatin plus ezetimibe, and 4620 to placebo. Allocation to simvastatin plus ezetimibe yielded an average LDL cholesterol difference of 33 mg/dL (SE 0.02, with about two-thirds of the sample adherent) during a median follow-up of 4.9 years, and produced a 17% proportional reduction in major atherosclerotic events (526 [11.3%] simvastatin plus ezetimibe vs. 619 [13.4%] placebo; rate ratio [RR] 0.83, 95% CI 0.74–0.94, log-rank \( p = 0.0021 \)). Patients allocated to simvastatin plus ezetimibe did not differ with respect to nonfatal myocardial infarction or death from coronary heart disease (213 [4.6%] vs. 230 [5.0%]; RR 0.92, 95% CI 0.76–1.11, \( p = 0.37 \)), and there were significant reductions in non-hemorrhagic stroke (131 [2.8%] vs. 174 [3.8%]; RR 0.75, 95% CI 0.60–0.94, \( p = 0.01 \)) and arterial revascularization procedures (284 [6.1%] vs. 352 [7.6%]; RR 0.79, 95% CI 0.68–0.93, \( p = 0.0036 \)). Adjustment for subgroup-specific reductions in LDL-c did not reveal evidence of differences between the proportional effects on major atherosclerotic events and the summary rate ratio in any subgroup examined, and, in particular, in patients on dialysis vs. those who were not on dialysis. The study concluded that reduction of LDL cholesterol with simvastatin 20 mg plus ezetimibe 10 mg daily safely reduced the incidence of major atherosclerotic events in patients with advanced chronic kidney disease [122].

- A sub-analysis of the treating to new targets study investigated how intensive lipid lowering with 80 mg of atorvastatin affects renal function when compared with 10 mg in patients with coronary heart disease. A total of 10,001 patients with coronary heart dis-

### Table 9 Recommendation for statin treatment according to cardiovascular risk category in diabetes

| Risk category       | Statin treatment |
|---------------------|------------------|
| LOW RISK            | Optional*        |
| INTERMEDIATE RISK   | Recommended      |
| HIGH RISK           | Highly recommended |
| VERY HIGH RISK      | Mandatory        |

* Optional means that non-pharmacological (lifestyle) measures are acceptable, provided that an LDL-c target <100 mg/dL is attained and maintained. For patients with LDL-c >160 mg/dL, statins are advisable at any risk category.
ease and LDL-c levels < 130 mg/dL were randomly assigned to double-blind therapy with 10 or 80 mg/d atorvastatin. Estimated GFR using the modification of diet in renal disease equation was compared at baseline and at the end of follow-up in 9656 participants with complete renal data. The expected 5-year decline in renal function was not observed. However, estimated GFR improved in both treatment groups, but was significantly greater with 80 mg than with 10 mg, suggesting this benefit may be dosage-related [123].

- Another sub-analysis of the TNT study investigated the effects of intensive lipid lowering with atorvastatin in patients with coronary heart disease (CHD) with and without preexisting chronic kidney disease (CKD). The study concluded that aggressive lipid lowering with atorvastatin 80 mg was both safe and effective in reducing excess cardiovascular events in a high-risk population with CKD and CHD [124].

42. In patients with diabetes and class III–IV heart failure, initiation of statin therapy is not recommended because there is no clear evidence of benefit in this group. [III, A]

Summary of evidence

- The effect of rosuvastatin in patients with chronic heart failure (GISSI-HF) randomized, multicenter clinical trial evaluated rosuvastatin 10 mg/day compared to placebo in 2285 patients with heart failure due to any cause or condition (New York Heart Association classes II–IV); 26% also had diabetes. There was no benefit in the outcomes of interest (death and hospitalization for cardiovascular causes) [125].

- The controlled rosuvastatin multinational trial in heart failure (CORONA) randomized study compared the use of rosuvastatin 10 mg versus placebo in 5011 patients aged >60 years with class II–IV heart failure of ischemic etiology (including 29% with diabetes). The primary endpoint was a composite of cardiovascular death, acute nonfatal MI, and nonfatal stroke during 36 months. Despite a 45% reduction in LDL-c, there was no significant between-group difference in the primary endpoint. The results were extensive to patients with diabetes in the subgroup analysis, due to low heterogeneity [126].

- A retrospective analysis of the CORONA trial compared 10 mg rosuvastatin daily with placebo in patients with ischemic systolic heart failure according to baseline high sensitivity-C reactive protein (hs-CRP) < 2.0 mg/L (placebo, n = 779; rosuvastatin, n = 777) or ≥ 2.0 mg/L (placebo, n = 1694; rosuvastatin, n = 1711). The primary outcome was cardiovascular death, myocardial infarction, or stroke. The study demonstrated a significant interaction between hs-CRP and the effect of rosuvastatin for most endpoints, whereby rosuvastatin treatment was associated with better outcomes in patients with hs-CRP ≥ 2.0 mg/L [127]. In addition, patients with heart failure due to ischemic heart disease who had NT-proBNP values < 103 pmol/L (868 pg/mL) had the best prognosis and, if assigned to rosuvastatin rather than placebo, had a greater reduction in the primary endpoint (HR 0.65, 95% CI 0.47–0.88) than patients in the other tertiles (heterogeneity test, p = 0.0192). This reflected fewer atherothrombotic events and sudden deaths in the active group, and may show a benefit from rosuvastatin use [128].

43. In the patient with diabetes and mild to moderate hypertriglyceridemia (TG 150–400 mg/dL), the combination of a statin and a fibrate is not usually recommended for reduction of cardiovascular risk. However, in the specific situation of a patient with triglycerides >204 mg/dL and HDL-c < 34 mg/dL, the combination of fenofibrate and a statin can be considered when lifestyle modifications have failed. [IIa, B]

Summary of evidence

- The pre-specified subgroup analysis of patients with diabetes from the ACCORD-LIPID (action to control cardiovascular risk in diabetes-lipids arm) study [129], comparing micronized fenofibrate 160 mg plus simvastatin 20–40 mg versus simvastatin 20–40 mg alone plus fenofibrate placebo, showed no reduction in the primary outcome. However, there was benefit in the pre-specified subgroup analysis of patients with triglycerides >204 mg/dL and HDL-c < 34 mg/dL.

- The FIELD (fenofibrate intervention and event lowering in diabetes) multinational RCT, randomized 9795 individuals with type 2 diabetes mellitus (aged 50–75 years, 2131 with previous cardiovascular disease and 7664 without) not on statin treatment at study enrollment to receive micronized fenofibrate 200 mg daily (n = 4895) or matching placebo (n = 4900) for 5 years of follow up. The primary outcome was coronary heart disease death or nonfatal myocardial infarction. The pre-specified outcome for subgroup analyses was total cardiovascular events (the composite of cardiovascular death, myocardial infarction, stroke, and coronary and carotid revascularization). Fenofibrate did not reduce risk of the primary outcome. However, it reduced the secondary pre-specified outcome of total cardiovascular events, due to fewer nonfatal myocardial infarctions and revascularizations [130].
Module 6. Management of hypertension

Targets

44. In patients with diabetes without clinical atherosclerotic disease (CLAD), blood pressure targets of a systolic blood pressure (SBP) < 130 mmHg and a diastolic blood pressure (DBP) < 80 mmHg may be reasonable, if well tolerated by the patient. [IIb, B]

Summary of evidence

- In the ACCORD study [131] of 4733 diabetic patients, randomization to an SBP target <120 mmHg vs. <140 mmHg could not reduce significantly the risk of the study’s primary outcome (HR 0.88, 95% CI 0.73–1.06, p = 0.20). Thus, the results of the study do not support the recommendations for stricter BP targets in this patient population. The mean SBP achieved in the first year of treatment in this trial were 119.3 mmHg for the <120 mmHg arm and 133.5 mmHg for the 140 mmHg arm, respectively. However, in the SBP < 120 mmHg arm, there was a 41% reduction in risk of stroke (HR 0.59, 95% CI 0.39–0.89, p = 0.01) with a low incidence of adverse events.

- The ACCORD BP study used a 2 × 2 factorial design, which also included comparisons of standard or intensive glycemic targets combined with intensive or standard blood pressure control in the same trial. A secondary pre-specified analysis [132] showed that, when combining intensive glycemic control with intensive blood pressure control, the rate of major CVD outcomes was significantly lowered when compared with combined standard BP and standard glycemic control.

- In a network meta-analysis including 42 clinical trials with random allocation into anti-hypertensive medication, control, or treatment target, a total of 144,220 individuals were compared in different strata of systolic blood pressure (SBP) to define the best target to reduce cardiovascular disease and all-cause mortality. In 30 trials, patients with type 2 diabetes were included. Patients were analyzed according to their mean achieved SBP in nine strata: 120–124; 125–129; 130–134; 135–139; 140–144; 145–149; 150–154; 154–159; and >160 mmHg. There were linear associations between mean achieved SBP and the risk of cardiovascular disease and mortality, with the lowest risk in the lowest stratum (120–124 mmHg). Individuals who achieved SBP 120–124 mmHg had a HR for all-cause mortality of 0.73 (95% CI 0.58–0.93) compared to those in the SBP 130–134 mmHg stratum: HR 0.59 (95% CI 0.45–0.77). Thus, reducing SBP levels to below 130 mmHg is associated with significant reductions in cardiovascular disease and in all-cause mortality [133].

45. In patients with established coronary heart disease (CLAD), it is not recommended to reduce blood pressure below 120/70 mmHg. [III, B]

Summary of evidence

- Because coronary perfusion occurs mainly during diastole, patients with coronary artery disease (CAD) could be at increased risk for coronary events if DBP falls below critical levels. A secondary analysis of data from the International Verapamil-Trandolapril Study (INVEST), including 22,576 patients with hypertension and CAD, determined whether low blood pressure could be associated with excess mortality and morbidity in this population. The analysis found a progressive increase for the risk for the primary outcome, all-cause death, and MI, but not stroke, with low DBP. The authors concluded that excessive reduction in diastolic pressure should be avoided in patients with CAD who are being treated for hypertension [134].

- Data from 22,672 patients with stable coronary artery disease from 45 countries enrolled in the CLARIFY registry and treated for hypertension were analyzed to ascertain whether a relationship exists between achieved blood pressure rates and cardiovascular events. SBP and DBP before each event were averaged and categorized into 10-mmHg increments. The primary outcome was a composite of cardiovascular death, MI, or stroke. Hazard ratios (HR) were estimated with multivariable adjusted Cox proportional hazards models, using 120–129 mmHg SBP and 70–79 mmHg DBP subgroups as references. The study concluded that, in patients with hypertension and coronary artery disease from routine clinical practice, SBP < 120 mmHg and DBP < 70 mmHg were each associated with adverse cardiovascular outcomes, including mortality, supporting the existence of a J-curve phenomenon. Thus, caution is advised in the use of antihypertensive treatment in patients with coronary artery disease [135].

- The ARIC (Atherosclerosis Risk In Communities) cohort of 11,565 adults analyzed associations between DBP and high-sensitivity cardiac troponin T (hs-cTnT) levels, as well as prospective associations between DBP and CV events. Compared with persons who had DBP 80–89 mmHg at baseline (ARIC visit 2), the adjusted odds ratio of having hs-cTnT ≥ 14 ng/L at that visit was 2.2 and 1.5 in those with DBP < 60 mmHg and 60–69 mmHg, respectively. Low DBP at baseline was also independently associated with progressive myocardial damage on the basis of estimated annual change in hs-cTnT over the 6 years between ARIC visits 2 and 4. In addition, compared with a DBP of 80–89 mm
Hg, a DBP < 60 mmHg was associated with incident CHD and mortality, but not with stroke. The DBP and incident CHD association was strongest with baseline hs-cTnT ≥ 14 ng/L (p value for interaction <0.001). Associations of low DBP with prevalent hs-cTnT and incident CHD were most pronounced among patients with baseline SBP ≥ 120 mmHg. The study concluded that, among adults with an SBP ≥ 120 mmHg (and, thus, elevated pulse pressure), low DBP was associated with subclinical myocardial damage and CHD events. When titrating treatment to SBP < 140 mmHg, it may be prudent to ensure that DBP levels do not fall below 70 mmHg and, particularly, not below 60 mmHg [136].

46. In patients with diabetes aged 80 years or older, a systolic blood pressure target <150 mmHg is reasonable. [IIa, B]

Summary of evidence
- In the hypertensive elderly (age ≥ 80 years), there is no evidence of benefits deriving from BP levels <140 mmHg, but there is an increased likelihood of adverse effects. The HYVET Study supports the recommendation of a BP target <150/90 mmHg, with a reduction in the risk of stroke and HF [137, 138]. The presence of isolated systolic hypertension (ISH) requires care regarding excessive reduction in DBP, which should be maintained over 60 mmHg or even over 65 mmHg in the presence of CAD [139].
- The SPRINT study reported a 24% reduction in the risk of the study’s primary outcome in elderly patients (age ≥75 years) allocated to the more intense BP treatment arm (mean SBP achieved, 123.4 mmHg) as compared to the group of standard SBP reduction (mean BP achieved, 134.8 mmHg). This occurred regardless of degree of frailty, with no increase in the number of adverse events in relation to the rest of the study population [140]. That suggests that BP targets for the elderly should be defined in the same as for other adults. It should be noted, however, that BP reduction should be performed carefully, considering comorbidities and the use of multiple medications.

47. In patients with stage III hypertension (defined as blood pressure ≥180/110 mmHg), the initial target blood pressure should be <140/90 mmHg. [I, A]

Summary of evidence
- In a meta-analysis, Thomopoulos et al. investigated if treatment to lower blood pressure benefits all grades of hypertension and determined the target BP levels to maximize outcome reduction. Significant outcome reductions were found independently of hypertension grade. No trend was observed toward changes in risk ratio with increasing baseline BP. In 32 RCTs (128,232 individuals), relative and absolute outcome reductions were significant for the SBP differences across 150 and 140 mmHg cutoffs. Below 130 mmHg, only stroke and all-cause mortality were significantly reduced. There was a significant trend toward greater absolute outcome reduction with lower SBP cutoffs. In 29 RCTs (107,665 individuals), outcomes were significantly reduced across DBP cutoffs of 90 and 80 mmHg. After excluding RCTs with baseline DBP <90 mmHg, only stroke reduction was significant at achieved DBP <80 mmHg. In conclusion, meta-analyses favor BP-lowering treatment in all grades of hypertension, at low-to-moderate risk, and lowering SBP/DBP to less than 140/90 mmHg. Achieving <130/80 mmHg appears safe, but only adds further reduction in stroke [141].

48. In patients with diabetes and increased albuminuria (>30 mg/g of creatinine), it is recommended that systolic blood pressure and diastolic blood pressure targets should be <130 and <80 mmHg respectively. [I, A]

Summary of evidence
- In the ADVANCE randomized clinical trial [142], 11,140 patients with type 2 diabetes and hypertension were randomized to receive with a fixed combination of perindopril and indapamide or matching placebo, in addition to current therapy. The primary endpoints were composites of major macrovascular and microvascular events (death from cardiovascular disease, nonfatal stroke, or nonfatal myocardial infarction) and new or worsening renal or diabetic eye disease. Analysis was by intention-to-treat. The macrovascular and microvascular composites were analyzed jointly and separately. Patients assigned to active therapy had a mean SBP reduction of 5.6 mmHg and a mean DBP reduction of 2.2 mmHg compared to the placebo arm. The relative risk of a major macrovascular or microvascular event was reduced by 9% (861 [15.5%] active vs. 938 [16.8%] placebo; HR 0.91, 95% CI 0.83–1.00, p = 0.04). The separate reductions in macrovascular and microvascular events were similar, but not independently significant (macrovascular: 0.92, 0.81–1.04, p = 0.16; microvascular: 0.91, 0.80–1.04, p = 0.16).
- In the IRMA-2 multinational, double-blind RCT, 590 hypertensive patients with type 2 diabetes and microalbuminuria were enrolled to receive irbesartan 150 mg daily or 300 mg daily for 2 years. The primary outcome was time to onset of diabetic nephropathy, defined by persistent albuminuria in overnight specimens, with a urinary albumin excretion rate >200 mcg/min and at least 30% higher than the
baseline level. Ten of the 194 patients in the 300-mg group (5.2%) and 19 of the 195 patients in the 150-mg group (9.7%) reached the primary endpoint, compared with 30 of the 201 patients in the placebo group (14.9%) (HR 0.30, 95% CI 0.14–0.61, p < 0.001, and HR 0.61, 95% CI 0.34–1.08, p = 0.081 for the two irbesartan groups, respectively). The average blood pressure during the course of the study was 144/83 mmHg in the placebo group, 143/83 mmHg in the 150-mg group, and 141/83 mmHg in the 300-mg group (p = 0.004 for the comparison of systolic blood pressure between the placebo group and the combined irbesartan groups) [143].

**Treatment**

49. The choice of initial drug therapy for hypertension should be based on efficacy, tolerability, cost, and presence of comorbidities. In general, diuretics, ACE inhibitors, angiotensin receptor blockers, or calcium channel blockers can be useful as initial monotherapy. [IIa, B]

**Summary of evidence**

- The randomized, double-blind, active-controlled antihypertensive and lipid-lowering treatment to prevent heart attack trial (ALLHAT), conducted from February 1994 through March 2002, evaluated 33,357 participants (age ≥ 55 years) with hypertension and at least one additional CHD risk factor from 623 North American centers to determine if calcium channel blockers or angiotensin-converting enzyme (ACE) inhibitors would lower the incidence of CHD or other CV events vs. treatment with a diuretic. The primary outcome was combined fatal CHD or non-fatal myocardial infarction, analyzed by intention to treat. Secondary outcomes were all-cause mortality, stroke, combined CHD (primary outcome, coronary revascularization, or angina with hospitalization), and combined CVD (combined CHD, stroke, treated angina without hospitalization, heart failure, and peripheral arterial disease). The primary outcome occurred in 2956 participants, with no difference between treatments. Compared with chlortalidone (6-year rate, 11.5%), the relative risks (RRs) were 0.98 (95% CI 0.90–1.07) for amlodipine (6-year rate, 11.3%) and 0.99 (95% CI 0.91–1.08) for lisinopril (6-year rate, 11.4%). Likewise, all-cause mortality did not differ between groups. For amlodipine vs. chlortalidone, secondary outcomes were similar except for a higher 6-year rate of heart failure with amlodipine (10.2% vs. 7.7%; RR, 1.38; 95% CI 1.25–1.52). For lisinopril vs. chlortalidone, lisinopril had higher 6-year rates of combined CVD (33.3% vs. 30.9%; RR, 1.10; 95% CI 1.05–1.16); stroke (6.3% vs. 5.6%; RR, 1.15; 95% CI 1.02–1.30); and HF (8.7% vs. 7.7%; RR, 1.19; 95% CI 1.07–1.31) [144].
- An analysis of the ALLHAT study to determine if treatment with a calcium channel blocker or an ACE inhibitor would decrease clinical complications compared with treatment with a thiazide-type diuretic in DM, IFG, and normoglycemia provided no evidence of superiority for treatment with calcium channel blockers or ACE inhibitors compared with a thiazide-type diuretic during first-step antihypertensive therapy in these populations [145].
- A meta-analysis of 354 randomized, double-blind, placebo-controlled trials of thiazides, beta blockers, ACE inhibitors, angiotensin II receptor antagonists, and calcium channel blockers in fixed dose was performed. Placebo adjusted reductions in systolic and diastolic blood pressure and prevalence of adverse effects, according to dose expressed as a multiple of the standard (recommended) doses of the drugs, were the main outcomes. All five classes produced similar reductions in blood pressure, with average SBP and DBP reductions of 9.1 and 5.5 mmHg respectively at standard doses and 7.1 and 4.4 mmHg respectively (20% lower) at half-standard doses. The drugs reduced blood pressure from all pretreatment levels, more so from higher levels; for a 10 mmHg-higher blood pressure, the reduction was 1.0 mmHg greater in SBP and 1.1 mmHg greater in DBP. The BP-lowering effects of different drug classes were additive. In addition, combination low-dose treatment increased efficacy and reduced adverse effects. From the average blood pressure in people who have strokes (150/90 mmHg), three drugs at half-standard dose were estimated to lower blood pressure by 20 mmHg systolic and 11 mmHg diastolic, thereby reducing the risk of stroke by 63% and the risk of ischemic heart disease events by 46% in the 60–69 age range [146].

50. In patients with diabetes and urinary albumin >30 mg/g, treatment with ACE inhibitors or angiotensin receptor blockers is indicated. [I, A]

**Summary of evidence**

- The reduction in end points in noninsulin-dependent diabetes mellitus with the angiotensin II antagonist losartan (RENAAL) study [147] investigated if albuminuria, a marker of renal disease, could also be a monitor of the renoprotective efficacy of RAS intervention by the angiotensin II (Ang II) antagonist losartan in patients with diabetic nephropathy. Data from the double-blind randomized RENAAL trial were used to examine the effects of losartan on a renal outcome (primary composite endpoint of doubling of serum
The MICRO-HOPE study investigated whether the use of irbesartan or amlodipine, or their combination, could lower the risk of cardiovascular endpoints in patients with type 2 diabetes.

The study concluded that cardiovascular benefit was observed regardless of the combination therapy used. The primary endpoint was a composite of cardiovascular death, myocardial infarction, or stroke, with a median follow-up of 4.5 years. The results showed a significant reduction in the risk of the primary endpoint (HR 0.77, 95% CI 0.64–0.92, p = 0.003) for the combination therapy compared to control.

The study also found that irbesartan was beneficial for cardiovascular outcomes, while amlodipine was beneficial for renal outcomes. The combination therapy provided a synergistic effect, leading to a more significant reduction in the risk of the primary endpoint.

51. When using more than one antihypertensive to achieve target blood pressure, it is reasonable to combine either an ACE inhibitor or an ARB with a dihydropyridine calcium channel blocker. [IIa, B]

Summary of evidence

- The ACCOMPLISH (Avoiding Cardiovascular Events Through COmbination Therapy in Patients Living With Systolic Hypertension) substudy [150], was designed to determine which combination therapy in patients with hypertension and diabetes most effectively decreased cardiovascular events. The outcomes effects of the ACE inhibitor benazepril, combined with amlodipine (B+A) or hydrochlorothiazide (B+H), were analyzed separately in diabetic patients as a pre-specified endpoint. A total of 6946 patients with diabetes were randomized to treatment with B+A or B+H. A subgroup of 2842 diabetic patients at very high risk (previous CV events or stroke) was also analyzed, as were 4559 patients without diabetes. The primary endpoint was a composite of cardiovascular death, myocardial infarction, stroke, hospitalization for angina, resuscitated arrest, and coronary revascularization. In the full diabetes group, the mean achieved blood pressures in the B+A and B+H groups were 131.5/72.6 and 132.7/73.7 mmHg respectively; over 30 months of follow-up, there were 207 (8.8%) and 287 (11.0%) primary events (HR 0.79, 95% CI 0.68–0.92, p = 0.003). For the diabetic patients at very high risk, there were 195 (13.6%) and 244 (17.3%) primary events (HR 0.77, 95% CI 0.64–0.93, p = 0.007). In the non-diabetic patients, there...
were 245 (10.8%) and 296 (12.9%) primary events (HR 0.82, 95% CI 0.69–0.97, p = 0.020). In the diabetic patients, B+A therapy had clear coronary benefits on both acute clinical events (p = 0.013) and revascularizations (p = 0.024). In patients with diabetes and hypertension, the calcium channel blocker amlodipine is superior to the diuretic hydrochlorothiazide when added to a renin-angiotensin system blocker for reduction of cardiovascular events in patients with diabetes requiring management of hypertension.

52. A combination of 3 or more drugs (ACE inhibitor or ARB plus amlodipine and a thiazide diuretic) can be useful in achieving BP goals. [IIa, B]

Summary of evidence
- A meta-analysis by Psaty [151] summarized the available clinical trial evidence concerning the safety and efficacy of various antihypertensive therapies used as first-line agents in terms of major cardiovascular disease endpoints and all-cause mortality. Network meta-analysis was used to combine direct within-trial between-drug comparisons with indirect evidence from other trials. Indirect comparisons preserving within-trial randomized findings were constructed from trials that had one treatment in common. Data were combined from 42 clinical trials that included 192,478 patients randomized to 7 major treatment strategies, including placebo. For all outcomes, low-dose diuretics were superior to placebo for coronary heart disease (CHD; RR 0.79, 95% CI 0.69–0.92); congestive heart failure (CHF; RR 0.51, 95% CI 0.42–0.62); stroke (RR 0.71, 95% CI 0.63–0.81); CV events (RR 0.76, 95% CI 0.69–0.83); CV mortality (RR 0.81, 95% CI 0.73–0.92); and total mortality (RR 0.90, 95% CI 0.84–0.96). None of the first-line treatment strategies—beta blockers, ACE inhibitors, calcium channel blockers (CCBs), alpha blockers, and angiotensin receptor blockers—was significantly better than low-dose diuretics for any outcome. Compared with CCBs, low-dose diuretics were associated with reduced risks of CV events (RR 0.94, 95% CI 0.89–1.00) and CHF (RR 0.74, 95% CI 0.67–0.81). Compared with ACE inhibitors, low-dose diuretics were associated with reduced risks of CHF (RR 0.88, 95% CI 0.80–0.96), CV events (RR 0.94, 95% CI 0.89–1.00), and stroke (RR 0.86, 0.77–0.97). Compared with beta blockers, low-dose diuretics were associated with a reduced risk of CV events (RR 0.89, 95% CI 0.80–0.98). Compared with alpha blockers, low-dose diuretics were associated with reduced risks of CHF (RR 0.51, 95% CI 0.43–0.60) and CV events (RR 0.84, 95% CI 0.75–0.93). Blood pressure changes were similar between comparison treatments. Low-dose diuretics were the most effective first-line treatment for preventing the occurrence of CV-related morbidity and mortality.

53. A combination of an ACE inhibitor and an ARB or a renin blocker is NOT recommended, due to the greater risk of loss of renal function, syncope, and hyperkalemia. [III, A]

Summary of evidence
- The ALTITUDE study asked whether the use of aliskiren would reduce cardiovascular and renal events in patients with type 2 diabetes and chronic kidney disease, cardiovascular disease, or both. The trial was stopped prematurely after the second interim efficacy analysis, because, after a median follow-up of 32.9 months, the primary endpoint (composite of the time to CV death or a first occurrence of cardiac arrest with resuscitation; nonfatal myocardial infarction; nonfatal stroke; unplanned hospitalization for heart failure; end-stage renal disease, death attributable to kidney failure, or need for renal replacement therapy with no dialysis or transplantation available or initiated; or doubling of the serum creatinine level from baseline) had occurred in 783 patients (18.3%) assigned to aliskiren as compared with 732 (17.1%) assigned to placebo (HR 1.08, 95% CI 0.98–1.20, p = 0.12). Thus, data do not support the addition of aliskiren to standard therapy with renin-angiotensin system blockade in patients with type 2 diabetes who are at high risk of cardiovascular and renal events. In fact, aliskiren may even be harmful [152].
- The ASTRONAUT study [153] was designed to investigate whether adding aliskiren to standard therapy would reduce the rate of CV death or readmission among HHF (hospitalization for heart failure) patients. Eligible patients were aged ≥18 years, with left ventricular ejection fraction (LVEF) 40% or less, elevated natriuretic peptides (brain natriuretic peptide [BNP] ≥ 400 pg/mL or N-terminal pro-BNP [NT-proBNP] ≥ 1600 pg/mL), and signs and symptoms of fluid overload. All patients received 150 mg of aliskiren (increased to 300 mg as tolerated) or placebo daily, in addition to standard therapy. The study drug was continued after discharge for a median 11.3 months. The main outcome measures were CV death or HF rehospitalization at 6 months and 12 months. In total, 1639 patients were randomized, with 1615 patients included in the final efficacy analysis cohort (808 aliskiren, 807 placebo). At randomization, patients were receiving diuretics (95.9%), beta blockers (82.5%), ACE inhibitors or ARBs (84.2%), and mineralocorticoid receptor antagonists (57.0%).
Summary of evidence

In total, 24.9% of patients receiving aliskiren (77 CV deaths, 153 HF readmissions) and 26.5% of patients receiving placebo (85 CV deaths, 166 HF readmissions) experienced the primary endpoint at 6 months (HR 0.92, 95% CI 0.76–1.12, p = 0.41). At 12 months, the event rates were 35.0% for the aliskiren group (126 CV deaths, 212 HF readmissions) and 37.3% for the placebo group (137 CV deaths, 224 HF readmissions; HR 0.93, 95% CI 0.79–1.09, p = 0.36). The rates of hyperkalemia, hypotension, and renal impairment/failure were higher in the aliskiren group than in the placebo arm. Among patients hospitalized for HF with reduced LVEF, initiation of aliskiren in addition to standard therapy did not reduce CV death or HF readmission at 6 months or 12 months after discharge.

- In patients who have vascular disease or high-risk diabetes without heart failure, ACE inhibitors reduce mortality and morbidity from cardiovascular causes, but the role of ARBs in such patients is unknown. The ACE inhibitor ramipril, the ARB telmisartan, and a combination of the two drugs in patients with vascular disease or high-risk diabetes were compared in the ONTARGET study [154]. A total of 8576 patients were assigned to receive 10 mg of ramipril per day, 8542 were assigned to receive 80 mg of telmisartan per day, and 8502 were assigned to receive both drugs. The primary composite outcome (death from cardiovascular causes, myocardial infarction, stroke, or hospitalization for heart failure) occurred in 16.5% of patients in the ramipril group, 16.7% in the telmisartan group, and 16.3% in the combined-therapy group (differences were not statistically significant). However, more adverse events were seen in patients randomized to combined therapy. In conclusion, the combination of the two drugs was associated with more adverse events, without increased benefit.

Module 7: Rationale for antiplatelet therapy

54. In patients with diabetes without clinical atherosclerotic disease (Table 5) in the HIGH-RISK category (Table 2), aged >65 years and with low risk of bleeding, acetylsalicylic acid can be useful. [IIa, B]

Summary of evidence

- The Japanese Primary Prevention of Atherosclerosis with Aspirin for Diabetes (JPAD) trial was designed to examine the efficacy of low-dose aspirin for primary prevention of atherosclerotic events in patients with type 2 diabetes and no previous cardiovascular events. The study randomized 1262 patients to receive aspirin (81 mg or 100 mg) and 1277 patients to a non-aspirin group. Mean (SD) age was 65 (10) years, and 55% were men; 58% of patients had hypertension, 53% had dyslipidemia, and BP and HbA1c were well controlled in both groups. The median follow-up period was 4.37 years and 193 patients were lost to follow-up, with data for those patients censored at the day of last follow-up. There was no reduction in the risk of CV events with low-dose aspirin for high-risk patients with diabetes in primary prevention. However, the event rate was lower than expected overall, and these findings should be interpreted in context with the low incidence of atherosclerotic disease in Japan and current management of cardiovascular risk factors [157].
- A meta-analysis of randomized controlled trials was performed to evaluate the benefits and harms of low-dose aspirin in people with diabetes and no cardiovascular disease. Six studies were eligible, with 10,117 participants. When aspirin was compared with placebo, there was no statistically significant reduction in the risk of major CV events (five studies, n = 9584; RR 0.90, 95% CI 0.81–1.00), CV mortality (four studies, n = 8557; RR 0.94, 95% CI 0.72–1.23), or all-cause mortality (four studies, n = 8557; RR 0.93, 95% CI 0.82–1.05). There was significant heterogeneity in analysis for myocardial infarction (I² = 62.2%; p = 0.02) and stroke (I² = 52.5%; p = 0.08). Aspirin significantly reduced the risk of myocardial infarction in men (0.57, 0.34–0.94) but not in women (1.08, 0.71–1.65; p value for interaction = 0.056). Evidence relating to harms was inconsistent [156].

55. In patients with diabetes without clinical atherosclerotic disease (Table 5) in the HIGH-RISK category (Table 2), aged >65 years and with low risk of bleeding, acetylsalicylic acid can be useful. [IIa, B]
SBP $\geq$ 140 mmHg and/or DBP $\geq$ 90 mmHg; “target attained” group, SBP < 140 mmHg and DBP < 90 mmHg) demonstrated that the incidence of the primary atherosclerotic events, especially cerebrovascular events, was higher in the unattained group than in the attained group. The incidence of cerebrovascular events was higher in the unattained group than in the attained group in patients without aspirin therapy; however, the incidence of cerebrovascular events in the unattained group was as low as the incidence in the attained group in patients on aspirin therapy. Cox proportional hazards analysis revealed that BP level was an independent predictor of cerebrovascular events in diabetic patients [158].

In a meta-analysis of RCTs with aspirin including 14 trials (107,686 participants), aspirin was associated with reductions in major cardiovascular events (risk ratio 0.90, 95% CI 0.85–0.95), myocardial infarction (risk ratio 0.86, 95% CI 0.75–0.93), ischemic stroke (risk ratio 0.86, 95% CI 0.75–0.98), and all-cause mortality (risk ratio 0.94, 95% CI 0.89–0.99). However, there were increases in hemorrhagic stroke (risk ratio 1.34, 95% CI 1.01–1.79) and major bleeding (risk ratio 1.55, 95% CI 1.35–1.78) with aspirin. The number needed to treat to prevent 1 major cardiovascular event over a mean follow-up of 6.8 years was 284. By comparison, the number needed to harm to cause 1 major bleeding was 299. In subgroup analyses, pooled results demonstrated a reduction in myocardial infarction among men (RR 0.71, 95% CI 0.59–0.85) and ischemic stroke among women (RR 0.77, 95% CI 0.63–0.93). Aspirin use was associated with a reduction (RR 0.65, 95% CI 0.51–0.82) in myocardial infarction among diabetic men. The results of meta-regression analyses suggested that aspirin therapy might be associated with a decrease in stroke among diabetic women and a decrease in MI among diabetic men, and that risk reductions achieved with low doses (75 mg/day) were as large as those obtained with higher doses (650 mg/day). The study concluded that low-dose aspirin was beneficial for primary prevention of CVD, and that the decision regarding an aspirin regimen should be made on an individual patient basis. The effects of aspirin therapy varied by sex and diabetes status [159].

56. In VERY HIGH-RISK patients, including those with clinical atherosclerotic disease (CLAD) and prior cardiovascular events (secondary prevention), antiplatelet therapy is indicated. [I, A]

**Summary of evidence**

- A collaborative meta-analysis of randomized trials of an antiplatelet regimen versus control or of one antiplatelet regimen versus another in high-risk patients (with acute or previous vascular disease or some other predisposing condition) was performed [6]. Trials had to use a randomization method that precluded prior knowledge of the next treatment to be allocated, and comparisons had to have study groups that differed only in terms of antiplatelet regimen. A total of 287 studies were included, involving 135,000 patients in comparisons of antiplatelet therapy versus control and 77,000 in comparisons of different antiplatelet regimens. Aspirin (or another oral antiplatelet drug) was protective in most types of patient at increased risk of occlusive vascular events, including those with AMI or ischemic stroke, unstable or stable angina, previous MI, stroke or cerebral ischemia, peripheral arterial disease, or atrial fibrillation [160].

57. In VERY HIGH-RISK patients with aspirin allergy or gastric intolerance, clopidogrel should be considered as an acceptable alternative. [IIa, B]

**Summary of evidence**

- The Clopidogrel vs. Aspirin in Patients at Risk of Ischemic Events (CAPRIE) trial was a randomized, blinded, multicenter trial of 19,185 patients with atherosclerotic disease manifested as recent ischemic stroke or myocardial infarction or symptomatic peripheral arterial disease. The number of readmissions for ischemic events (defined as angina, transient ischemic attack, or limb ischemia) or bleeding events was determined for the entire cohort. A significant reduction in the total number of readmissions for ischemic events or bleeding was seen with clopidogrel use compared with aspirin (1502 vs. 1673, $p = 0.010$) over an average of 1.6 years of treatment. This reduction in rehospitalization was consistent across individual outcomes of angina, transient ischemic attack, or limb ischemia, or bleeding. Clopidogrel also resulted in a 7.9% relative risk reduction in a combined endpoint of vascular death, stroke, myocardial infarction, or rehospitalization for ischemic events or bleeding (15.1–13.7% at 1 year, $p = 0.011$) as compared with aspirin [161].

58. Dual antiplatelet therapy is recommended for at least 1 year in VERY HIGH-RISK patients after acute coronary syndrome. [I, A]

**Summary of evidence**

- The TRITON-TIMI 38 trial aimed to randomly compare prasugrel (a new thienopyridine antiplatelet agent) vs. clopidogrel in 13,608 patients with moderate-to-high-risk acute coronary syndromes scheduled to undergo percutaneous coronary intervention. Prasugrel was given in a 60-mg loading dose and a 10-mg daily maintenance dose, while clopidogrel was
In the CHARISMA trial, 15,603 patients with either non-ST-elevation acute coronary syndrome undergoing PCI were randomly assigned to double-blind treatment with clopidogrel \((n = 1313)\) or placebo \((n = 1345)\). Patients were pretreated with aspirin and clopidogrel for a median of 6 days before PCI during the initial hospital admission, and for a median of 10 days overall. After PCI, most patients \((>80\%)\) in both groups received open-label thienopyridine for about 4 weeks, after which clopidogrel was restarted for a mean of 8 months. The primary endpoint was a composite of cardiovascular death, myocardial infarction, or urgent target-vessel revascularization within 30 days of PCI, in an intention-to-treat analysis. Fifty-nine \((4.5\%)\) patients in the clopidogrel group reached the primary endpoint, compared with 86 \((6.4\%)\) in the placebo group \((RR 0.70, 95\% CI 0.50–0.97, p = 0.03)\). Long-term administration of clopidogrel after PCI was associated with a lower rate of CV death, myocardial infarction, or any revascularization \((p = 0.03)\) and of CV death or myocardial infarction \((p = 0.047)\). Overall, including events before and after PCI, there was a 31\% reduction in CV death or myocardial infarction \((p = 0.002)\), and, at follow-up, there was no significant difference in major bleeding between the groups \((p = 0.64)\) [163].

In the CHARISMA trial, 15,603 patients with either clinically evident cardiovascular disease or multiple risk factors were randomized to receive clopidogrel \((75 \text{ mg per day})\) plus low-dose aspirin \((75–162 \text{ mg per day})\) or placebo plus low-dose aspirin and followed for a median of 28 months. The primary endpoint was a composite of myocardial infarction, stroke, or death from cardiovascular causes. The rate of the primary efficacy endpoint was 6.8\% with clopidogrel plus aspirin and 7.3\% with placebo plus aspirin \((RR 0.93, 95\% CI 0.83–1.05, p = 0.22)\). The principal secondary efficacy endpoint, which included hospitalizations for ischemic events, occurred in 16.7\% and 17.9\% \((RR 0.92, 95\% CI 0.86–0.995, p = 0.04)\), and the rate of severe bleeding was 1.7\% and 1.3\% \((RR 1.25, 95\% CI 0.97–1.61, p = 0.09)\). In patients with multiple risk factors, the rate of the primary endpoint was 6.6\% with clopidogrel and 5.5\% with placebo \((RR 1.2, 95\% CI 0.91–1.59, p = 0.20)\), and the rate of death from cardiovascular causes was also higher with clopidogrel \((3.9\% \text{ vs. } 2.2\%, p = 0.01)\). In subgroup analysis of patients with clinically evident atherothrombosis, the rate was 6.9\% with clopidogrel and 7.9\% with placebo \((RR 0.88, 95\% CI 0.77–0.998, p = 0.046)\), suggesting benefit with clopidogrel treatment in patients with symptomatic atherothrombosis and harm in patients with multiple risk factors. Overall, clopidogrel plus aspirin was not significantly more effective than aspirin alone in reducing the rate of myocardial infarction, stroke, or death from cardiovascular causes [164].

In the multicenter, double-blind, randomized PLATO trial, ticagrelor \((180-\text{mg loading dose, } 90 \text{ mg twice daily thereafter})\) and clopidogrel \((300–600-\text{mg loading dose, } 75 \text{ mg daily thereafter})\) were compared for the prevention of CV events in 18,624 patients admitted to hospital with an acute coronary syndrome, with or without ST-segment elevation. At 12 months, the primary composite endpoint \((death \text{ from vascular causes, myocardial infarction, or stroke})\) had occurred in 9.8\% of patients receiving ticagrelor as compared with 11.7\% of those receiving clopidogrel \((p < 0.001)\). The rate of death from any cause was also reduced with ticagrelor \((4.5\% \text{ vs. } 5.9\% \text{ with clopidogrel}, p < 0.001)\). No significant difference in rates of major bleeding was found between the ticagrelor and clopidogrel groups \((p = 0.43)\), but ticagrelor was associated with a higher rate of major bleeding not related to coronary-artery bypass grafting \((4.5\% \text{ vs. } 3.8\%, p = 0.03)\), including more instances of fatal intracranial bleeding and fewer of fatal bleeding of other types [165].

The PEGASUS study investigated the efficacy and safety of ticagrelor after an acute coronary syndrome, in a double-blind 1:1:1 fashion. The trial randomized 21,162 patients who had had a myocardial infarction 1–3 years earlier to ticagrelor 90 mg twice daily, ticagrelor 60 mg twice daily, or placebo. All patients received low-dose aspirin and were followed for a
median of 33 months. The primary efficacy endpoint was the composite of cardiovascular death, myocardial infarction, or stroke. The primary safety endpoint was Thrombolysis in Myocardial Infarction (TIMI)-defined major bleeding. Both ticagrelor doses reduced the rate of the primary efficacy endpoint, with Kaplan–Meier rates at 3 years of 7.85% in the 90-mg ticagrelor group, 7.77% in the 60-mg ticagrelor group, and 9.04% in the placebo group (HR for 90-mg ticagrelor vs. placebo: 0.85, 95% CI 0.75–0.96, \( p = 0.008 \); HR for 60-mg ticagrelor vs. placebo: 0.84, 95% CI 0.74–0.95, \( p = 0.004 \)). Rates of TIMI major bleeding were higher with ticagrelor (2.60% with 90 mg and 2.30% with 60 mg) than with placebo (1.06% (\( p < 0.001 \)) for each dose vs. placebo); the rates of intracranial hemorrhage or fatal bleeding in the three groups were 0.63, 0.71, and 0.60% respectively. Therefore, in patients with a previous myocardial infarction, at least one year earlier, treatment with ticagrelor significantly reduced the risk of CV death, myocardial infarction, and stroke, but increased the risk of major bleeding [166].

59. In patients who are not at high risk of bleeding complications, continuation of dual antiplatelet therapy may be reasonable for longer than 12 months after acute coronary syndrome. [IIb, A]

Summary of evidence

- In the PEGASUS study, the efficacy and safety of ticagrelor after an acute coronary syndrome was investigated in a double-blind 1:1:1 fashion. The trial randomized 21,162 patients who had had a myocardial infarction 1–3 years earlier to ticagrelor at a dose of 90 mg twice daily, ticagrelor at a dose of 60 mg twice daily, or placebo. All the patients received low-dose aspirin and were followed for a median of 33 months. The primary efficacy endpoint was the composite of cardiovascular death, myocardial infarction, or stroke. The primary safety endpoint was Thrombolysis in Myocardial Infarction (TIMI) major bleeding. Ticagrelor in both doses reduced the rate of the primary efficacy endpoint, with Kaplan–Meier rates at 3 years of 7.85% in the group receiving 90 mg of ticagrelor twice daily, 7.77% in the group receiving 60 mg of ticagrelor twice daily, and 9.04% in the placebo group (hazard ratio for 90 mg of ticagrelor vs. placebo, 0.85; 95% CI 0.75–0.96; \( p = 0.008 \); hazard ratio for 60 mg of ticagrelor vs. placebo, 0.84; 95% CI 0.74–0.95; \( p = 0.004 \)). Rates of TIMI major bleeding were higher with ticagrelor (2.60% with 90 mg and 2.30% with 60 mg) than with placebo (1.06%) (\( p < 0.001 \)) for each dose vs. placebo); the rates of intracranial hemorrhage or fatal bleeding in the three groups were 0.63, 0.71, and 0.60% respectively. Therefore, in patients with a previous myocardial infarction, at least one year earlier, treatment with ticagrelor significantly reduced the risk of cardiovascular death, myocardial infarction, or stroke, and increased the risk of major bleeding [166]. The DAPT study sought to investigate if 30 months of DAPT was superior to 12 months in patients undergoing DES and bare-metal stent (BMS) PCI. A total of 9961 patients were randomized at 452 sites in 11 countries: 5020 to prolonged DAPT and 4941 to placebo. Approximately 30% had diabetes mellitus, 25% were smokers and 6% had peripheral arterial disease. Patients were enrolled 72 h after stent placement and were given open-label aspirin and thienopyridine for 12 months, per current practice norms. Indication for PCI was stable angina in 38%, ST-segment elevation myocardial infarction (STEMI) in 10% and NSTE-acute coronary syndrome (NSTE-ACS) in 32%. Approximately two-thirds of the patients received clopidogrel, whereas the rest received prasugrel. At 12 months, patients without an ischemic or bleeding complication and with documented compliance, were randomized in a 1:1 fashion to receive an additional 18 months of DAPT or matching placebo. The primary endpoint of major adverse cardiac and cerebrovascular events (MACCE) was significantly lower in the continued DAPT arm compared with placebo (4.3% vs. 5.9%, hazard ratio 0.71, 95% confidence interval 0.59–0.85, \( p < 0.001 \)). There were reductions in all MI (2.1% vs. 4.1%, \( p < 0.001 \)) and stent thrombosis (0.4% vs. 1.4%, \( p < 0.001 \)), but all-cause mortality was higher (2.0% vs. 1.5%, \( p = 0.05 \)), driven mostly by an increase in non-cardiovascular deaths (1% vs. 0.5%, \( p = 0.002 \)), including cancer-related death (0.62% vs. 0.28%, \( p = 0.02 \)) and bleeding-related death (0.22% vs. 0.06%, \( p = 0.06 \)). Moderate and severe GUSTO bleeding was also higher with DAPT (2.5% vs. 1.6%, \( p = 0.001 \)), as was BARC 2, 3, or 5 bleeding (5.6% vs. 2.9%, \( p < 0.001 \)). The DAPT study showed that longer duration of DAPT following PCI results in lower stent thrombosis and recurrent MIs, but higher bleeding and all-cause mortality compared with a 12-month duration [167].

Conclusions

Although cardiovascular risk is increased in patients with diabetes when compared to age-matched nondiabetic individuals, recent evidence indicates that there is a high prevalence of lower-risk individuals among this population. Risk stratification is clearly needed, either to intensify more effective preventive measures in high-risk categories or to avoid overtreatment of lower-risk
patients. The present Panel structured a risk-based guide to help clinicians optimize cardiovascular prevention in diabetes. The Panel recovered the concept of treating-to-target, as they are considered important to promote better adherence to treatment and can be useful for clinicians to improve prevention in clinical practice. In the present guideline, there is a clear shift toward a more intensive treatment in the very-high risk category, especially regarding lipid-lowering therapy with statins, where new, lower lipid targets are proposed. The Panel understands that patients with diabetes at very high risk have very high mortality and one of the most important currently available actions to reduce residual risk is to obtain further reductions in LDL-c levels. The panel also reviews the potential role of the new anti-hyperglycemic drugs in reducing cardiovascular risk, as well as hypertension targets and drug choice. Finally, we also propose a practical guideline to guide decision-making about screening for silent coronary artery disease. We understand that intensifying treatment may increase costs to the health care system; however, the number of avoided events and lives saved clearly outweighs these costs. The Brazilian Diabetes Society, the Brazilian Cardiology Society, and the Brazilian Endocrinology Society are now united in the task to reduce cardiovascular disease in patients with diabetes.

Abbreviations

4D: die deutsche diabetes dialyse; ABI: ankle-brachial index; ACCOMPLISH: Avoiding Cardiovascular Events Through ComBination Therapy in Patients Living With Systolic Hypertension; ACCORD: action to control cardiovascular risk in diabetes; ACCORD-LIPID: action to control cardiovascular risk in diabetes-lipids arm; ACE: angiotensin-converting enzyme; ADVANCE: action in diabetes and vascular disease: preterax and diamicon modified release controlled evaluation; AMI: acute myocardial infarction; ARIC: atherosclerosis risk in communities; AUC-ROC: area under the ROC curve; AURORA: a study to evaluate the use of rosuvastatin in subjects on regular hemodialysis; BMS: bare-metal stent; CAC: coronary artery calcium; CAD: coronary artery disease; CAPRIE: clopidogrel vs. aspirin in patients at risk of ischemic events; CARDIA: coronary artery risk development in young adults; CARD: Collaborative Atorvastatin Diabetes Study; CBBS: calcium channel blockers; CCTA: coronary computed tomography angiography; CHD: coronary heart disease; CHF: congestive heart failure; CIMT: carotid-artery intima-media thickness; CKD: chronic kidney disease; CLAD: clinical atherosclerotic disease; CMRI: cardiac magnetic resonance imaging; CORONA: controlled rosuvastatin multinational trial in heart failure; pCP: carotid plaque burden; CV: cardiovascular; CVD: cardiovascular disease; DCCT: diabetes control and complications trial; DIAD: detection of ischemia in asymptomatic diabetics; DYNAMIT: do you need to assess myocardial ischemia in type-2 diabetes; ECG: electrocardiogram; EDIC: epidemiology of diabetes interventions and complications; ESRD: end-stage renal disease; FT: familiar hypercholesterolemia; FIELD: fenofibrate intervention and event lowering in diabetes; GISSI-HF: effect of rosvastatin in patients with chronic heart failure; HHF: hospitalization for heart failure; HR: hazard ratios; ICU: intensive care unit; INVEST: International Verapamil-Trandolapril Study; ISh: isolated systolic hypertension; JPAD: Japanese Primary Prevention of Atherosclerosis with Aspirin for Diabetes; LVEF: left ventricular ejection fraction; MACCE: major adverse cardiac and cerebrovascular events; MACE: major adverse cardiovascular events; MI: myocardial infarction; MPS: myocardial perfusion imaging; NRI: net reclassification index; NSTEMI: non-ST-segment elevation myocardial infarction; PCI: percutaneous coronary intervention; PCI-SS: subclinical atherosclerosis; RAS: renin-angiotensin system; RR: rate ratio; RRR: relative risk reduction; SCAT: subclinical atherosclerosis; SMI: silent myocardial ischemia; SPECT: single-photon emission computed tomography; ST: ST-segment elevation myocardial infarction; TACT: treat to new targets; UKPDS: United Kingdom Prospective Diabetes Study; VADT: veterans affairs diabetes trial.

Authors’ contributions

All authors had full participation in the search for references, in the development of contents, and in drafting the text. MCB acted as the lead author. All authors read and approved the final manuscript.

Author details

1 Departamento de Medicina Interna, Faculdade de Medicina, Universidade Federal do Rio Grande do Sul (UFRGS), Rua Ramiro Barcelos, 2400, Porto Alegre, RS 90035-003, Brazil. 2 Serviço de Medicina Interna, Hospital de Clínicas de Porto Alegre (HCIP), UFRGS, Rua Ramiro Barcelos, 2350, Porto Alegre, RS 90035-903, Brazil. 3 Instituto Estadual de Diabetes e Endocrinologia Luiz Capriglione, Rua Moncorvo Filho, 90, Rio de Janeiro, RJ 20211-340, Brazil. 4 Faculdade de Medicina de Valença (FMV), Rua Sebastião Dantas Moreira, 40, Valença, RJ 27600-000, Brazil. 5 Faculdade de Medicina da Universidade Presidente Antônio Carlos (FAME/UNIPAC), Av. Juiz de Fora, 1100, Juiz De Fora, MG 36048-000, Brazil. 6 Instituto Dante Pazzanese de Cardiologia, Av. Dante Pazzanese, 500, São Paulo, SP 04012-180, Brazil. 7 Universidade Federal de São Paulo (UNIFESP), Rua Loeuffen, 1350, São Paulo, SP 04040-001, Brazil. 8 UFRGS, Rua Ramiro Barcelos, 2350, Porto Alegre, RS 90035-903, Brazil. 9 Universidade de São Paulo (USP), Av. Dr. Enéas de Carvalho Aguiar, 44, São Paulo, SP 05403-000, Brazil. 10 Faculdade de Ciências Médicas de Minas Gerais, Alameda Ezequiel Dias, 275, Belo Horizonte, MG 30130-110, Brazil. 11 Universidade Federal do Goiás (UFG), 14ª Avenida, s/n, Setor Leste Universitário, Goiânia, GO 74605-020, Brazil. 12 Pontifícia Universidade Católica de Campinas (PUC-Campinas), Av. John Boyd Dunlop, s/n, Campinas, SP 13059-900, Brazil. 13 Faculdade de Ciências, Médicas da Santa Casa de São Paulo, Rua Dr. Cesário Motta Jr, 112, São Paulo, SP 01221-020, Brazil. 14 Universidade Federal de Santa Catarina (UFSC), Rua Profa. Maria Flora Pausewang, s/n, Florianópolis, SC 88040-970, Brazil. 15 Clínica de Endocrinologia e Metabologia, Av. Tancredo Neves, 1632/708, Salvador, BA 41820-020, Brazil. 16 Centro de Diabetes de Belo Horizonte, Rua Niquel, 31, Belo Horizonte, MG 30220-280, Brazil. 17 Faculdade de Medicina de São José do Rio Preto, Av. Brg. Faria Lima, 5416, São José do Rio Preto, SP 15090-000, Brazil. 18 UNIFESP, Rua Botucatu, 740, São Paulo, SP 04023-002, Brazil. 19 UNIFESP, Rua Botucatu, 740, São Paulo, SP 04023-002, Brazil.

Acknowledgements

Not applicable.

Competing interests

ROM has received speaker honorarium from: Novartis Pharmaceuticals, Novo Nordisk, Merck Serono, Sanofi-Aventis, Ache, and Eli Lilly. CMV has received honoraria as speaker from Novo Nordisk, Takeda, Novartis, and Aegerion Pharmaceuticals.

SV over the last 5 years, has received honoraria for clinical research from Amgen/Novartis/Novo Nordisk/Bayer/EMS/Boehringer Ingelheim/Cristália and Sanofi-Aventis: Advisory Board to Novo Nordisk/Novartis and Boehringer Ingelheim; has received honoraria as speaker from Novo Nordisk/Novartis and Boehringer Ingelheim.

FT has received honoraria for medical lectures from: Ache, AstraZeneca, Boehringer Lilly, Lilly, Novo Nordisk, Sanofi, Servier, and Takeda.

MHN participated in the Trial Assessing Long Term Use of PCSK9 Inhibition in Subjects With Genetic LDL Disorders (TAUSSIG), sponsored by Amgen. He also participated in events sponsored by MSD, Sanofi/Regeneron, Amgen, and was sponsored by Sanofi for participation in conferences.

RDS over the last 3 years has received honoraria for consulting, research and speaker activities from: Agenon, Aegerion, AstraZeneca, Akcea, Biolab, Boehringer Ingelheim, Cerenis, Genzyme, Eli Lilly, Merck, Sanofi/Regeneron, Pfizer, Procaps, Torrent, Unilever, and Kowa.

The other authors have no competing interests to declare.

Availability of data and materials

Data sharing is not applicable to this article as no datasets were generated or analyzed during the current study.
References

1. Bertoluci MC, Pimazoni-Netto A, Pires AC, Pessaro AE, Schaan BD, Caramelli B, Polanczyk CA, Junior CV, Guallandro DM, Malerbi DA, et al. Diabetes and cardiovascular disease: from evidence to clinical practice—position statement 2014 of Brazilian Diabetes Society. Diabetol Metab Syndr. 2014;6:58.

2. Bertoluci M, Rocha VZ. Cardiovascular risk assessment in patients with diabetes. Diabetol Metab Syndr. 2017;9:1–13.

3. Booth GL, Kapral MK, Fung K, Tu JV. Relation between age and cardiovascular disease in men and women with diabetes compared with non-diabetic people: a population-based retrospective cohort study. Lancet. 2006;368:29–36.

4. Wannamethee SG, Shaper AG, Whincup PH, Lennon L, Sattar N. Impact of diabetes on cardiovascular disease risk and all-cause mortality in older men: influence of age at onset, diabetes duration, and established and novel risk factors. Arch Intern Med. 2011;171:404–10.

5. Li R, O’Sullivan MJ, Robinson J, Safford MM, Curb D, Johnson KC. Family history of myocardial infarction predicts incident coronary heart disease in postmenopausal women with diabetes: the Women’s Health Initiative Observational Study. Diabetes Metab Res Rev. 2009;25:725–32.

6. Mottillo S, Filion KB, Genest J, Joseph L, Pilote L, Poirier P, Rinfore S, Schiffin EL, Eisenberg MJ. The metabolic syndrome and cardiovascular risk: a systematic review and meta-analysis. J Am Coll Cardiol. 2010;56:1113–22.

7. James PA, Oparil S, Carter BL, Cushman WC, Dennison-Himmelfarb C, Handler J, Lackland DT, LeFevre ML, MacKenzie TD, Ogedegbe O, et al. 2014 evidence-based guideline for the management of high blood pressure in adults: report from the panel members appointed to the Eighth Joint National Committee (JNC 8). JAMA. 2014;311:507–20.

8. Njolstad I, Arnesen E, Lund-Larsen PG. Smoking, serum lipids, blood pressure, and sex differences in myocardial infarction. A 12-year follow-up of the Finnmark Study. Circulation. 1996;93:450–60.

9. Chronic Kidney Disease Prognosis Consortium, Matsushita K, van der Velde M, Astor BC, Woodward M, Levey AS, de Jong PE, Coreh J, Gansevoort RT. Association of estimated glomerular filtration rate and albuminuria with all-cause and cardiovascular mortality in general population cohorts: a collaborative meta-analysis. Lancet. 2016;375:2073–81.

10. Gerstein HC, Mann JF, Yi Q, Zinman B, Dinneen SF, Hoogeveen R, Halle JP, Young J, Rashkow A, Joyce C, et al. Albuminuria and risk of cardiovascular events, death, and heart failure in diabetic and non-diabetic individuals. JAMA. 2001;286:421–6.

11. Pop-Busui R, Evans GW, Gerstein HC, Fonseca V, Fleg JL, Hoogeveen R, Gennuth S, Grunus RH, Conson MA, Prineas R, et al. Effects of cardiac autonomic dysfunction on mortality risk in the action to control cardiovascular risk in diabetes (ACCORD) trial. Diabetes Care. 2010;33:1578–84.

12. Gerstein HC, Ambrosius WT, Danis R, Irsay-Beigi F, Cushman W, Calles J, Banerji M, Schubart U, Chew EL, Group AS. Diabetic retinopathy, its progression, and incident cardiovascular events in the ACCORD trial. Diabetes Care. 2013;36:1266–71.

13. Kramer CK, Rodrigues TC, Canani LH, Gross JL, Azevedo MJ. Diabetic retinopathy predicts all-cause mortality and cardiovascular events in both type 1 and 2 diabetes: meta-analysis of observational studies. Diabetes Care. 2011;34:1238–44.

14. Akazawa S, Toyikubo M, Nakano Y, Nakamura S, Tamai H, Yonemoto K, Sadasima E, Kawasaki T, Koga N. Usefulness of carotid plaque (sum and maximum of plaque thickness) in combination with intima-media thickness for the detection of coronary artery disease in asymptomatic patients with diabetes. J Diabetes Investig. 2016;7:396–403.

15. Min JK, Labounty TM, Gomez MJ, Achenbach S, Al-Mallah M, Budoff MJ, Cademartini F, Callister TQ, Chang MJ, Cheng V, et al. Incremental prognostic value of coronary computed tomographic angiography over coronary artery calcium score for risk prediction of major adverse cardiac events in asymptomatic diabetic individuals. Atherosclerosis. 2014;232:298–304.

16. Li J, Luo Y, Xu Y, Yang J, Zheng L, Hasimbu BY, Yu J, Hu D. Risk factors of peripheral arterial disease and relationship between low ankle—brachial index and mortality from all-cause and cardiovascular disease in Chinese patients with type 2 diabetes. Cinc J. 2007;71:377–81.

17. Bown MJ, Sutton AJ, Bell PR, Sayers RD. A meta-analysis of 50 years of ruptured abdominal aortic aneurysm repair. Br J Surg. 2002;89:714–30.

18. Glimaker H, Holmberg L, Elvin A, Nybacka O, Almgren B, Bjorck CG, Eriksson I. Natural history of patients with abdominal aortic aneurysm. Eur J Vasc Surg. 1991;5:125–30.

19. Semmens JB, Norman PE, Lawrence-Brown MM, Holman CD. Influence of gender on outcome from ruptured abdominal aortic aneurysm. Br J Surg. 2000;87:191–4.

20. Lederle FA, Johnson GR, Wilson SE, Aneurysm D. Management veterans affairs cooperative S. Abdominal aortic aneurysm in women. J Vasc Surg. 2001;34:122–6.

21. Bath MF, Gokani VJ, Sidloff DA, Jones LR, Choke E, Sayers RD, Bown MJ. Systematic review of cardiovascular disease and cardiovascular death in patients with a small abdominal aortic aneurysm. Br J Surg. 2015;102:866–72.

22. Lieber A, Jorgens J. Cinefluorography of coronary artery calcification. Correlation with clinical arteriosclerotic heart disease and autopsy findings. Am J Roentgenol Radium Ther Nucl Med. 1961;86:1063–72.

23. DeTrano R, Guerci AD, Carr JJ, Bild DE, Burke G, Folsom AR, Liu K, Shea S, Skolz M, Bluemke DA, et al. Coronary calcium as a predictor of coronary events in four racial or ethnic groups. N Engl J Med. 2008;359:1336–45.

24. Erbel R, Mohlenkamp S, Moebus S, Schmermund A, Lehnmann N, Stang A, Dragano N, Gronehagem B, Seibel R, Kalsch H, et al. Coronary risk stratification, discrimination, and reclassification improvement based on quantification of subclinical coronary atherosclerosis: the Heinz Nixdorf Recall study. J Am Coll Cardiol. 2010;56:1397–406.

25. Rahmani S, Nakanishi R, Budoff MJ. Imaging atherosclerosis in diabetes: current state. Curr Diab Rep. 2016;16:105.

26. Raggi P, Shaw LJ, Berman DS, Callister TQ. Prognostic value of coronary artery calcium screening in subjects with and without diabetes. J Am Coll Cardiol. 2004;43:1663–9.

27. Elkeles RS, Godland IF, Feher MD, Rubens MB, Roughton M, Nugara F, Humphries SE, Richmond W, Flather MD, Group PS. Coronary calcium measurement improves prediction of cardiovascular events in asymptomatic patients with type 2 diabetes: the PREDICT study. Eur Heart J. 2008;29:2244–51.

28. Silverman MG, Blaha MJ, Budoff MJ, Rivera JR, Raggi P, Shaw LJ, Berman D, Callister T, Rumberger JA, Rana JS, et al. Potential implications of coronary artery calcium testing for guiding aspirin use among asymptomatic individuals with diabetes. J Card Diab. 2012;35:624–6.

29. Carr JJ, Jacobs DR Jr, Terry RG, Shay CM, Sidney S, Liu K, Scheiner PJ, Lewis CE, Shikany JM, Reis JP, et al. Association of coronary artery calcium in adults aged 32 to 46 years with incident coronary heart disease and death. JAMA Cardiol. 2017;2:391.

30. Yehoja J, Young R, McClelland RL, Delaney JC, Polonsky TS, Dawood FZ, Blaha MJ, Miedema MD, Sibley CT, Carr JJ, et al. Utility of nontraditional risk markers in atherosclerotic cardiovascular disease risk assessment. J Am Coll Cardiol. 2016;67:139–47.

31. Kramer CK, Zinman B, Gross JL, Canani LH, Rodrigues TC, Azvedo MJ, Retnakaran R. Coronary artery calcium score prediction of all-cause mortality and cardiovascular events in people with type 2 diabetes: systematic review and meta-analysis. BMJ. 2013;346:f1654.

32. Agarwal S, Morgan T, Harringon DM, Xu J, Cox AJ, Freedman BI, Carr JJ, Bowden DW. Coronary calcium score and prediction of all-cause mortality in diabetes: the diabetes heart study. Diabetes Care. 2011;34:1219–24.
33. Agarwal S, Cox AJ, Herrington DM, Jorgensen NW, Xu J, Freedman BI, Carr JJ, Bowden DW. Coronary calcium score predicts cardiovascular mortality in diabetes: diabetes heart study. Diabetes Care. 2011;34:6972–7.

34. Nambi V, Chambless L, Folsom AR, He M, Hu Y, Mosley T, Volcik K, Boerwinkle E, Ballentine CM. Cardiac intima-media thickness and presence or absence of plaque improves prediction of coronary heart disease risk: the ARIC (Atherosclerosis Risk In Communities) study. J Am Coll Cardiol. 2010;55:1600–7.

35. Soliman EZ, Backlund JC, Bebu I, Orchard TJ, Zinman B, Lachin JM, Group DER. Electrocardiographic abnormalities and cardiovascular disease risk in type 1 diabetes: The Epidemiology of Diabetes Interventions and Complications (EDIC) Study. Diabetes Care. 2017;40:709.

36. Davis TM, Coleman RL, Holman RR, Group U. Prognostic significance of silent myocardial infarction in newly diagnosed type 2 diabetes mellitus: United Kingdom Prospective Diabetes Study (UKPDS) 79. Circulation. 2013;127:980–7.

37. Milan Study on Atherosclerosis and Diabetes (MiSAD) Group. Prevalence of unrecognized silent myocardial ischemia and its association with atherosclerotic risk factors in noninsulin-dependent diabetes mellitus. Am J Cardiol. 1997;79:134–9.

38. Young LH, Wackers FJ, Chyun DA, Davey JA, Barretti RJ, Taillefer R, Heller GV, Iskandrian AE, Wittlin SD, Filippich N, et al. Cardiac outcomes after screening for asymptomatic coronary artery disease in patients with type 2 diabetes: the DIAD study: a randomized controlled trial. JAMA. 2009;301:1547–55.

39. Lieve MM, Moulin P, Thivolet C, Rodier M, Righelau V, Fenfrosis A, Pradignac A, Ouvze M, Investigators D. Detection of silent myocardial ischemia in asymptomatic patients with diabetes: results of a randomized trial and meta-analysis assessing the effectiveness of systematic screening. Trials. 2011;12:23.

40. Zellweger MJ, Maram M, Osterheus HH, Keller U, Muller-Brand J, Jeger EF, Iskandrian AE, Wittlin SD, Filippich N, et al. Cardiac outcomes after screening for asymptomatic coronary artery disease in patients with type 2 diabetes: the DIAD study: a randomized controlled trial. JAMA. 2009;301:1547–55.

41. Lieve MM, Moulin P, Thivolet C, Rodier M, Righelau V, Fenfrosis A, Pradignac A, Ouvze M, Investigators D. Detection of silent myocardial ischemia in asymptomatic patients with diabetes: results of a randomized trial and meta-analysis assessing the effectiveness of systematic screening. Trials. 2011;12:23.

42. Zellweger MJ, Maram M, Osterheus HH, Keller U, Muller-Brand J, Jeger EF, Iskandrian AE, Wittlin SD, Filippich N, et al. Cardiac outcomes after screening for asymptomatic coronary artery disease in patients with type 2 diabetes: the DIAD study: a randomized controlled trial. JAMA. 2009;301:1547–55.

43. Anand DV, Lim E, Hopkins D, Corder R, Shaw LJ, Brunn AM, Gaultier-Bourgeois S, Bassand JP, Bernard Y. Use of dobutamine stress echocardiography responses and coronary heart disease mortality among men with diabetes mellitus. Circulation. 2008;117:2734–42.

44. Anand DV, Lim E, Hopkins D, Corder R, Shaw LJ, Sharp P, Lipkin D, Lahiri AK. Risk stratification in uncomplicated type 2 diabetes: prospective evaluation of the combined use of coronary artery calcium imaging and selective myocardial perfusion scintigraphy. Eur Heart J. 2006;27:131–21.

45. Fenfrosis A, Zimmermann C, Bouard D, Sabbah A, Menoueau N, Gaultier-Bourgeois S, Bassand JP, Bernard Y. Use of dobutamine stress echocardiography in detecting silent myocardial ischaemia in asymptomatic diabetic patients: a comparison with thallium scintigraphy and exercise testing. Diabet Med. 2001;18:900–5.

46. Greenwood JF, Nareda N, Younger JF, Brown JM, Nixon J, Everett CC, Bjerstvedt P, Ridgway JP, Radjenovic A, Richardson C, et al. Cardiovacular magnetic resonance imaging and selective myocardial perfusion scintigraphy for diagnosis of coronary heart disease (CE-MARC): a prospective trial. Lancet. 2012;379:453–60.

47. Muhlstein JB, Lappe DL, Lima JA, Rosen BD, May HT, Knight S, Bluemke DA, Towner SR, Le V, Bair TL, et al. Effect of screening for coronary artery disease using CT angiography on mortality and cardiac events in high-risk patients with diabetes: the FACTOR-64 randomized clinical trial. JAMA. 2014;312:2324–43.

48. Hadamitzky M, Hein F, Meyer T, Bischoff B, Martinoff S, Schomig A, Hausleiter J. Prognostic value of coronary computed tomographic angiography in diabetic patients without known coronary artery disease. Diabetes Care. 2010;33:1358–63.

49. The Diabetes Control and Complications Trial Research Group. The effect of intensive treatment of diabetes on the development and progression of long-term complications in insulin-dependent diabetes mellitus. N Engl J Med. 1993;329:977–86.

50. UK Prospective Diabetes Study (UKPDS) Group. Intensive blood-glucose control with sulphonylureas or insulin compared with conventional treatment and risk of complications in patients with type 2 diabetes (UKPDS 33). Lancet. 1998;352:837–53.

51. Nathan DM, Cleary PA, Backlund JY, Gennuth SM, Lachin JM, Orchard TJ, Raskin P, Zinman B. Intensive diabetes treatment and cardiovascular disease in patients with type 1 diabetes. N Engl J Med. 2005;353:2643–53.

52. Orchard TJ, Nathan DM, Zinman B, Cleary P, Brillou D, Backlund JY, Lachin JM. Association between 7 years of intensive treatment of type 1 diabetes and long-term mortality. JAMA. 2015;313:45–53.

53. Holman RR, Paul SK, Bethel MA, Matthews DR, Neil HA. 10-year follow-up of intensive glucose control in type 2 diabetes. N Engl J Med. 2008;358:2545–59.

54. Patel A, MacMahon S, Chalmers J, Neil B, Billot L, Woodward M, Marre M, Cooper M, Glasspo M, Grobbee DE, et al. Intensive blood glucose control and vascular outcomes in patients with type 2 diabetes. N Engl J Med. 2005;358:2560–72.

55. Duckworth W, Abraica C, Mortiz T, Reda D, Emanuele N, Reaven PD, Zieve F, Marks J, Davis SN, Hayward R, et al. Glucose control and vascular complications in veterans with type 2 diabetes. N Engl J Med. 2009;360:129–39.

56. Pogach L, Aron D. The other side of quality improvement in diabetes for seniors: a proposal for an overtreatment glycemiyic measure. Arch Intern Med. 2012;172:1510–2.

57. Lipska KJ, Krumholz H, Soones T, Lee SJ. Polypyrhmacology in the aging patient: a review of glycemic control in older adults with type 2 diabetes. JAMA. 2016;315:1034–45.

58. Holstein A, Patzer OM, Machalke K, Holstein JD, Stumvoll M, Kovacs P. Substantial increase in incidence of severe hypoglycemia between 1997–2000 and 2007–2010: a German longitudinal population based study. Diabetes Care. 2012;35:972–5.

59. Geller AI, Shehab N, Lovegrove MC, Kegler SR, Weidenbach K, Ryan GI, Budnitz DS. National estimates of insulin-related hypoglycemia and errors leading to emergency department visits and hospitalizations. JAMA Intern Med. 2014;174:678–86.

60. Lipska KJ, Inzucchi SE. Effect of glucose management on coronary heart disease risk in patients with diabetes. In: McGuire DK, Marx N, editors. Diabetes in cardiovascular disease: a companion to Braunwald’s heart disease. Philadelphia: Elsevier; 2015. p. 155–70.

61. Schaan BD, de Figueiredo Neto JA, Moreira LB, Ledur P, Mattos LAP, Magioni D, Precoma DB, Machado CA, Brasilieiro ALS, Pena FM, et al. Diabetes and cardiovascular events in high-risk patients: insights from a multicenter registry in a middle-income country. Diabetes Res Clin Pract. 2017;127:275–84.

62. Lipska KJ, Warton EM, Huang ES, Moffet HH, Inzucchi SE, Krumholz HM, Karter AJ. HbA1c and risk of severe hypoglycemia in type 2 diabetes: the Diabetes and Aging Study. Diabetes Care. 2013;36:5355–42.

63. Lipska KJ, Krumholz HM. Comparing diabetes medications: where do we set the bar? JAMA Intern Med. 2014;174:317–8.

64. Kirsh SR, Aron DC. Choosing targets for glycaemia, blood pressure and low-density lipoprotein cholesterol in elderly individuals with diabetes mellitus. Drugs Aging. 2011;28:945–60.

65. Malmberg K. Prospective randomised study of intensive insulin treatment on long term survival after acute myocardial infarction in patients with diabetes mellitus. DiGAMI (Diabetes Mellitus, Insulin Glucose Infusion in Acute Myocardial Infarction) Study Group. BMJ. 1997;314:1512–5.

66. Malmberg K, Ryden L, Wedel H, Birkeland K, Bootsm a A, Dickstein K, Efendic S, Fisher M, Hamsten A, Herlitz J, et al. Intense metabolic control by means of insulin in patients with diabetes mellitus and acute myocardial infarction (DIGAMI 2): effects on mortality and morbidity. Eur Heart J. 2005;26:650–61.
67. Cheung NW, Wong VW, McLean M. The Hyperglycemia: intensive Insulin Infusion in Infarction (HI-S) study: a randomized controlled trial of insulin infusion therapy for myocardial infarction. Diabetes Care. 2006;29:765–70.

68. Ledur P, Almeida L, Pellanda LC, Schaan BD. Clinical features and outcomes in patients with diabetes mellitus undergoing coronary artery bypass graft in a reference center in southern Brazil. Rev Assoc Med Bras. 1992;38(5):200–4.

69. Jones KW, Cain AS, Mitchell JH, Millar RC, Rimmasch HL, French TK, Abbate SL, Roberts CA, Stevenson SR, Marshall D, et al. Hyperglycemia predicts mortality after CABG: postoperative hyperglycemia predicts dramatic increases in mortality after coronary artery bypass graft surgery. J Diabetes Complicat. 2008;22:365–70.

70. Fornary AP, Zerr KJ, Grunkemeier GL, Starr A. Continuous intravenous insulin infusion reduces the incidence of deep sternal wound infection in diabetic patients after cardiac surgical procedures. Ann Thorac Surg. 1999;67:352–60 (discussion 60-2).

71. Fornary AP, Gao G, Grunkemeier GL, Wu Y, Zerr KJ, Bookin SQ, Floten HS, Starr A. Continuous insulin infusion reduces mortality in patients with diabetes undergoing coronary artery bypass grafting. J Thorac Cardiovasc Surg. 2003;125:1007–21.

72. van den Berghe G, Wouters P, Weekers F, Verwaest C, Bruyninckx F, Schetz M, Vlasselaers D, Ferdinande P, Lauwers P, Bouillon R. Intensive insulin therapy in critically ill patients. N Engl J Med. 2001;345:1359–67.

73. Finnis S, Chittock DR, Su SY, Blair D, Foster D, Dhingra V, Bellomo R, Cook D, Dodek P, Henderson WR, et al. Intensive versus conventional glucose control in critically ill patients. N Engl J Med. 2009;360:1283–97.

74. Griesdale DE, de Souza RV, van Dam RM, Heyland DK, Cook DJ, Dhillon A, Dhillon R, Henderson WR, Chittock DR, Finfer S, et al. Intensive insulin therapy and mortality among critically ill patients: a meta-analysis including NICE-SUGAR study data. CMAJ. 2009;180:821–7.

75. Lazar HL, McDonnell MM, Chipkin S, Fitzgerald C, Bliss C, Cabral H. Effects of aggressive versus moderate glycemic control on clinical outcomes in diabetic coronary artery bypass graft patients. Ann Surg. 2011;254:458–63 (discussion 63-4).

76. Kwon S, Thompson R, Dellinger P, Yanez D, Fanohki E, Flum D. Importance of peripereoperative glycemic control in general surgery: a report from the Surgical Care and Outcomes Assessment Program. Ann Surg. 2013;257:8–14.

77. Noordzij PG, Boersma E, Schreiner F, Kertai MD, Feringa HH, Dunkelgrun LC, Rodrigues TC, Azevedo MJ. Effect of antihyperglycemic agents added to metformin and a sulfonylurea on glycemic control and weight gain in type 2 diabetes: a network meta-analysis. Ann Intern Med. 2011;154:672–9.

78. Investigators OT, Gerstein HC, Bosch J, Dagenais GR, Diaz R, Jung H, Maggioni AP, Pogue J, Probstfield J, Ramachandran A, et al. Basal insulin and cardiovascular and other outcomes in dysglycemia. N Engl J Med. 2012;367:319–28.

79. Marso SP, McGuire DK, Zinman B, Poulter NR, Emerson SS, Reber TR, Pratley RE, Haahr PM, Lange M, Brown-Frandsen K, et al. Efficacy and safety of degludec versus glargine in type 2 diabetes. N Engl J Med. 2017. doi:10.1056/NEJMoa1615692.

80. Zinman B, Wariner C, Lachin JM, Fitchett D, Bluhmki E, Hantel S, Matthews M, Devins T, Johansen OE, Wöerle HJ, et al. Empagliflozin, cardiovascular outcomes, and mortality in type 2 diabetes. N Engl J Med. 2015;373:2117–28.

81. Neal B, Perkovic V, Mahaffey KW, de Zeeuw D, Fulcher G, Endretu N, Shaw W, Law G, Desai M, Matthews DR. Canagliflozin and cardiovascular and renal events in type 2 diabetes. N Engl J Med. 2017. doi:10.1056/NEJMoa1611925.

82. Marso SP, Daniels GH, Brown-Frandsen K, Kristensen P, Mann JF, Nauck MA, Nissen SE, Pocock S, Poulter NR, Ravn LS, et al. Liraglutide and cardiovascular outcomes in type 2 diabetes. N Engl J Med. 2016;375:311–22.

83. Marso SP, Bain SC, Consoli A, Elaschewitz FG, Jodar E, Leiter LA, Lingvay I, Rosenstock J, Seufert J, Warren ML, et al. Semaglutide and cardiovascular outcomes in patients with type 2 diabetes. N Engl J Med. 2016;375:1834–44.

84. Dormandy JA, Charbonnel B, Eckland DJ, Erdmann E, Massi-Benedetti M, Moulies IK, Skene AM, Tan MH, Lefebvre PJ, Murray GD, et al. Secondary prevention of macrovascular events in patients with type 2 diabetes in the PROactive Study (PROspective pioglitAzone Clinical Trial in Macrovascular Events): a randomised controlled trial. Lancet. 2005;366:1279–89.

85. Green JB, Bethel MA, Armstrong PW, Buse JB, Engel SS, Garg J, Josse R, Kaufman KD, Koglin J, Korn S, et al. Effect of sitagliptin on cardiovascular outcomes in type 2 diabetes. N Engl J Med. 2013;373:232–42.

86. Scicna BM, Bhatt DL, Braunwald E, Steg PG, Davidson J, Hirshberg B, Ohman P, Frederich R, Wiviott SD, Hoffman EB, et al. Saxagliptin and cardiovascular outcomes in patients with type 2 diabetes mellitus. N Engl J Med. 2013;369:1317–26.

87. White WB, Cannon CP, Heller SR, Nissen SE, Bergleren RM, Bakris GL, Perez AT, Fleck PR, Mehta CR, Kupfer S, et al. Alogliptin after acute coronary syndrome in patients with type 2 diabetes. N Engl J Med. 2013;369:1327–35.

88. Pfeffer MA, Claggett B, Diaz R, Dickstein K, Gerstein HC, Kober LV, Lawson FC, Ping L, Wei X, Lewis EF, et al. Lixisenatide in patients with type 2 diabetes and acute coronary syndrome. N Engl J Med. 2015;373:2247–57.

89. Rados DV, Pinto LC, Remonti LR, Leitao CB, Gross JL. Correction: the association between sulfonylurea use and all-cause and cardiovascular mortality: a meta-analysis with trial sequential analysis of randomized clinical trials. PLoS Med. 2016;13:e1002091.

90. Sabatine MS, Giugliano RP, Keesch AC, Honarpour N, Wiviott SD, Murphy SA, Kuder JF, Wang H, Liu T, Wasserman SM, Sever PS, Pedersen PR, FOURIER Steering Comitee and Investigators. Evolocumab and clinical outcomes in patients with cardiovascular disease. N Engl J Med. 2017;376(18):1713–22.

91. Cannon CP, Blazing MA, Giugliano RP, Maggioni A, White JA, Theroux P, Darius H, Lewis BS, Ophuis TO, Jukema JM, et al. Ezetimibe added
to statin therapy after acute coronary syndromes. N Engl J Med. 2015;372:2387–97.

101. Kearney PM, Blackwell L, Collins R, Keech A, Simes J, Petro R, Armittage J, Baigent C. Efficacy of cholesterol-lowering therapy in 18,686 people with diabetes: a 13 randomised trials of statins: a meta-analysis. Lancet. 2008;371:117–25.

102. Boekholdt SM, Hovingh GK, Mora S, Arsenault BJ, Amarenco P, Catapano AL, Cooney MT, Corra U, Cosyns B, Deaton C, et al. 2016 European guidelines on cardiovascular disease prevention in clinical practice: the Sixth Joint Task Force of the European Society of Cardiology and Other Societies on Cardiovascular Disease Prevention in Clinical Practice (constituted by representatives of 10 societies and by invited experts) Developed with the special contribution of the European Association for Cardiovascular Prevention & Rehabilitation (EACPR). Eur Heart J. 2016;37:2315–31.

103. Shepherd J, Barter P, Carmera R, Deedwania P, Fruchart JC, Haffner S, Hsia J, Breazna A, LaRosa J, Grundy S, et al. Efficacy of lowering LDL cholesterol substantially below currently recommended levels in patients with coronary heart disease and diabetes: the treating to new targets (TNT) study. Diabetes Care. 2006;29:1220–6.

104. Baigent C, Blackwell L, Emberson J, Holland LE, Reith C, Bhala N, Peto R, Blom DJ, Hala T, Bolognese M, Lillestol MJ, Toth PD, Burgess L, Sabatine MS, Giugliano RP, Wiviott SD, Raal FJ, Blom DJ, Robinson J, Ramasamy I. Recent advances in physiological lipoprotein metabolism. Zhang XL, Zhu QQ, Zhu L, Chen JZ, Chen QH, Li GN, Xie J, Kang LN, Xu Verbeek R, Stoekenbroek RM, Hovingh GK. PCSK9 inhibitors: novel approaches to statin therapy after acute coronary syndromes. N Engl J Med. 2015;371:117–25.

106. Colhoun HM, Ginsberg HN, Leiter LA, Chaudhari U, Lorenzato C, Pordy Colhoun HM, Betteridge DJ, Durrington PN, Hitman GA, Neil HA, Collins R, Armitage J, Parish S, Sleigh P, Peto R. MRC/BHF Heart Protection Study of cholesterol-lowering with simvastatin in 5963 people with diabetes: a randomised placebo-controlled trial. Lancet. 2003;361:2005–16.

107. Bedis US, Singh M, Singh PP, Bhunya R, Bahekar A, Molnar J, Khosla S, Arora R. Effects of statins on progression of carotid atherosclerosis as measured by carotid intimal–medial thickness: a meta-analysis of randomized controlled trials. J Cardiovasc Pharmacol Ther. 2010;15:268–73.

108. Filipi MC, Hoes AW, Agewall S, Albus C, Brotons C, Catapano AL, Cooney MT, Corra U, Cosyns B, Deaton C, et al. 2016 European guidelines on cardiovascular disease prevention in clinical practice: the Sixth Joint Task Force of the European Society of Cardiology and Other Societies on Cardiovascular Disease Prevention in Clinical Practice (constituted by representatives of 10 societies and by invited experts) Developed with the special contribution of the European Association for Cardiovascular Prevention & Rehabilitation (EACPR). Eur Heart J. 2016;37:2315–31.

109. Colhoun HM, Betteridge DJ, Durrington PN, Hitman GA, Neil HA, Collins R, Armitage J, Parish S, Sleigh P, Peto R. MRC/BHF Heart Protection Study of cholesterol-lowering with simvastatin in 5963 people with diabetes: a randomised placebo-controlled trial. Lancet. 2003;361:2005–16.
of combined cardiovascular risk factor management strategies in type 2 diabetes: the ACCORD randomized trial. Diabetes Care. 2014;37:1271–82.

133. Bundy JD, Li C, Stuchlik P, Bu X, Kelly TN, Mills KT, He H, Chen J, Whelton PK, He J. Systolic blood pressure reduction and risk of cardiovascular disease and mortality: a systematic review and network meta-analysis. JAMA Cardiol. 2017. doi:10.1001/jamacardio.2017.1421.

134. Messerli FH, Mancia G, Conti CR, Hewkin AC, Kupfer S, Champion A, Koloch R, Benetos A, Pepine CJ. Dogma disputed: can aggressively lowering blood pressure in hypertensive patients with coronary artery disease be dangerous? Ann Intern Med. 2006;144:884–93.

135. Vidal-Petiot E, Ford I, Greenlaw N, Ferrari R, Fox KM, Tardif JC, Tendera M, McEvoy JW, Chen Y, Rawlings A, Hoogeveen RC, Ballantyne CM, Hulley SB, Furberg CD, Gurland B, McDonald R, Perry HM, Schnaper HW, Wright JT Jr, Williamson JD, Whelton PK, Snyder JK, Sink KM, Rocco MV, Thomopoulos C, Parati G, Zanchetti A. Effects of blood pressure lowering with ramipril on cardiovascular and microvascular outcomes in people with diabetes mellitus: results of the HOPE study and MICRO-HOPE substudy. Lancet. 2000;355:253–9.

136. Weber MA, Bakris GL, Jamerson K, Weir M, Kjeldsen SE, Devereaux RB, Vélezazquez Ej, Dahirof B, Kelly RY, Hua TA, et al. Cardiovascular events during differing hypertension therapies in patients with diabetes. J Am Coll Cardiol. 2010;56:77–85.

137. Psaty BM, Lumley T, Furberg CD, Shellenbaum G, Pahor M, Alderman MH, Weiss NS. Health outcomes associated with various antihypertensive therapies used as first-line agents: a network meta-analysis. JAMA. 2003;289:2534–44.

138. Parving HH, Brenner BM, McMurray JJ, de Zeeuw D, Haefner SM, Solomon SD, Chaturvedi N, Persson F, Desai AS, Nicolaides M, et al. Cardioenal end points in a trial of aliskiren for type 2 diabetes. N Engl J Med. 2012;367:2204–13.

139. Gheorghide M, Bohm M, Greene SJ, Fonarow GC, Lewis EF, Zannad F, Solomon SD, Bascheria F, Botha J, Hua TA, et al. Effect of aliskiren on cardiovascular and heart failure readmissions among patients hospitalized for heart failure: the ASTRONAUT randomized trial. JAMA. 2013;309:1125–35.

140. Yusuf S, Teo KK, Pogue J, Dyal L, Copland I, Schumacher H, Dagenais G, Sleight P, Anderson C, Telomisartan, ramipril, or both in patients with high risk for vascular events: N Engl J Med. 2008;358:1547–59.

141. Antplatelet Trialists' Collaboration. Collaborative overview of randomised trials of antiplatelet therapy—I: prevention of death, myocardial infarction, and stroke by prolonged antiplatelet therapy in various categories of patients. BMJ. 1994;308:81–106.

142. De Berardis G, Sacco M, Strippoli GF, Graziano G, Tognoni G, Nicolucci A. Aspirin for primary prevention of cardiovascular events in people with diabetes: meta-analysis of randomised controlled trials. BMJ. 2009;339:b453.

143. Ogawa H, Nakayama M, Morimoto T, Uemura S, Kanachi M, Doi N, Inouchi H, Sugiyama S, Saito Y. Low-dose aspirin for primary prevention of atherosclerotic events in patients with type 2 diabetes: a randomized controlled trial. JAMA. 2008;300:2134–41.

144. Soejima H, Ogawa H, Morimoto T, Nakayama M, Okada S, Uemura S, Kanachi M, Doi N, Sakuma M, Inouchi H, et al. Aspirin reduces cerebrovascular events in type 2 diabetic patients with poorly controlled blood pressure. Subanalysis from the JPAD trial. Circ J. 2012;76:1526–32.

145. Xie M, Shan Z, Zhang Y, Chen S, Yang W, Bao W, Rong Y, Yu X, Hu FB, Liu L. Aspirin for primary prevention of cardiovascular events: meta-analysis of randomized controlled trials and subgroup analysis by sex and diabetes status. PLOS ONE. 2014;9:e90306.

146. Antithrombotic Trialists' Collaboration. Collaborative meta-analysis of randomised trials of antithrombotic therapy for prevention of death, myocardial infarction, and stroke in high risk patients. BMJ. 2002;324:71–86.

147. Bhatt DL, Hirsh AT, Ringleb PA, Hacke W, Topol EJ. Reduction in the need for hospitalization for recurrent ischemic events and bleeding with clopidogrel instead of aspirin: CAPRIE investigators. Am Heart J. 2000;140:67–73.

148. Atkins RC, Briganti EM, Lewis JB, Hunsicker LG, Braden G, Champion de Crespygien PJ, DeFierra G, Drury P, Locatelli F, Wiegmann TB, et al. Proteinuria reduction and progression to renal failure in patients with type 2 diabetes mellitus and overt nephropathy. Am J Kidney Dis. 2005;45:281–7.
165. Wallentin L, Becker RC, Budaj A, Cannon CP, Emanuelsson H, Held C, Horrow J, Husted S, James S, Katus H, et al. Ticagrelor versus clopidogrel in patients with acute coronary syndromes. N Engl J Med. 2009;361:1045–57.

166. Bonaca MP, Bhatt DL, Cohen M, Steg PG, Storey RF, Jensen EC, Magnani G, Bansilal S, Fish MP, Im K, et al. Long-term use of ticagrelor in patients with prior myocardial infarction. N Engl J Med. 2015;372:1791–800.

167. Mauri L, Kereiakes DJ, Yeh RW, Driscoll-Shempp P, Cutlip DE, Steg PG, Normand SL, Braunwald E, Wiviott SD, Cohen DJ, et al. Twelve or 30 months of dual antiplatelet therapy after drug-eluting stents. N Engl J Med. 2014;371:2155–66.