Amlodipine - a third generation dihydropyridine calcium antagonist

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H.-M. Steffen

Amlodipine, a third generation dihydropyridine calcium antagonist, is characterized by a higher vascular selectivity and a smaller negative inotropic effect compared to nifedipine. With its long elimination half-life and low variability in trough-to-peak plasma concentrations, once-daily application is possible without loss of therapeutic efficacy. Placebo-controlled and comparative studies with a variety of antianginal and antihypertensive agents have confirmed the efficacy of amlodipine in patients with arterial hypertension and/or coronary artery disease. As a result of gradual onset of action, amlodipine demonstrates no clinically significant stimulation of neuroendocrine systems and preliminary results have shown that it may be useful in patients with heart failure. Amlodipine is well tolerated and exerts no adverse or unfavourable effects on carbohydrate and lipid metabolism. As a result of these factors, amlodipine represents a therapeutic advance in the treatment of hypertension and coronary artery disease. J Clin Basic Cardiol 1999; 2: 45–52.

Key words: amlodipine, dihydropyridine, calcium antagonist, coronary artery disease, arterial hypertension, left ventricular hypertrophy, diabetes mellitus

Calcium antagonists (CAs) were first introduced into pharmacotherapy over 25 years ago as coronary vasodilators for the treatment of coronary heart disease and have since achieved notable recognition in the treatment of arterial hypertension [1]. The initial investigations of the working group with Albrecht Fleckenstein [2] led to the finding that the common effect of this very heterogeneous class of substances can be ascribed to a specific impairment of calcium ion influx through so-called slow-channels embedded in the cell membrane. In accordance with their chemical structure, CAs can be divided into four groups [3, 4], of which only the first three are important for cardiovascular therapy: phenylalkylamines (prototype: verapamil), dihydropyridines (DHPs, prototype: nifedipine), benzothiazepines (prototype: diltiazem), and diphenylalkylamines (prototype: cinnarizine).

On the basis of their biophysical and pharmacological properties, the four types of potential-dependent, calcium selective membrane channels found in many body tissues are differentiated and described as L- (long-term activation), T- (temporary opening), N- (neural), and P-type (Purkinje-cells). Conventional calcium antagonists act in this process as selective blockers of the L-type calcium channel which consists of five subdivisions described as α1, α2, β, γ, and δ with a molecular weight of approximately 400 kDa. The α1-subtype is the actual membrane pore and has specific binding sites for phenylalkylamines, DHPs, and benzothiazepines [5]. Mibebradil, a benzimidazolyl-tetraline represented the first T-type calcium channel blocker (CCB), but last year this drug was withdrawn worldwide from the market due to unforeseen drug interactions via the cytochrome P450 system.

Efforts to obtain substances with a higher tissue selectivity, longer duration of action and, compared to the parent substance, a less marked negative inotropy have led to the development of a new generation of CAs, in particular within the class of DHPs (see Table 1). Publications during the last 5 years have initiated a still ongoing debate about the safety of CAs, especially the DHPs [6–8], despite the fact that the efficacy and safety of nitrendipine has been demonstrated in the high risk group of elderly patients with systolic hypertension [9]. Also, a recent case-control study has shown that patients on long-acting CAs had no increased risk of cardiovascular events compared with those on β-blocker monotherapy [10], probably because of a lack of persistent stimulation of the sympathetic nervous system [11]. A retrospective cohort analysis of more than 1400 newly diagnosed hypertensive patients without coronary heart disease (CHD) also showed no increased relative risk for those treated with predominantly long-acting CCBs compared to β-blockers or diuretics [12]. Amlodipine, due to its unique pharmacokinetic and pharmacodynamic properties, is of major importance among the novel DHPs and will be described in greater detail hereafter.

Preclinical pharmacology

As a function of plasma concentration, amlodipine inhibits the contraction of vascular smooth muscle in depolarized rat aorta at a rate about double that of nifedipine; the maximum efficacy is achieved after three hours compared to 30 minutes for nifedipine [13]. In the guinea pig, concentrations required for a 50 % contraction inhibition are 160 times higher for amlodipine than for coronary arteries [5]. Amlodipine in a concentration of 100 nMol weakens the maximum contraction of vascular smooth muscle in depolarized rat aorta at a rate about double that of nifedipine; the maximum efficacy is achieved after three hours compared to 30 minutes for nifedipine [13]. In the guinea pig, concentrations required for a 50 % contraction inhibition are 160 times higher for amlodipine than for coronary arteries [5]. Amlodipine in a concentration of 100 nMol weakens the maximum contraction of vascular smooth muscle in depolarized rat aorta at a rate about double that of nifedipine; the maximum efficacy is achieved after three hours compared to 30 minutes for nifedipine [13]. In the guinea pig, concentrations required for a 50 % contraction inhibition are 160 times higher for amlodipine than for coronary arteries [5]. Amlodipine in a concentration of 100 nMol weakens the maximum contraction of vascular smooth muscle in depolarized rat aorta at a rate about double that of nifedipine; the maximum efficacy is achieved after three hours compared to 30 minutes for nifedipine [13]. In the guinea pig, concentrations required for a 50 % contraction inhibition are 160 times higher for amlodipine than for coronary arteries [5]. Amlodipine in a concentration of 100 nMol weakens the maximum contraction of vascular smooth muscle in depolarized rat aorta at a rate about double that of nifedipine; the maximum efficacy is achieved after three hours compared to 30 minutes for nifedipine [13]. In the guinea pig, concentrations required for a 50 % contraction inhibition are 160 times higher for amlodipine than for coronary arteries [5]. Amlodipine in a concentration of 100 nMol weakens the maximum contraction of arterial vessels > peripheral vessels

Cautions and contraindications

- Cautions: Patients with hepatic cirrhosis, severe renal insufficiency, and patients undergoing hemodialysis require reduced doses.
- Contraindications: Patients with aortic stenosis, severe congestive heart failure, severe left ventricular dysfunction, and patients with a history of hypersensitivity to amlodipine should be avoided.

Table 1. Selectivity and pharmacological properties of various calcium antagonists (modified in accordance with [5, 126, 127])

<table>
<thead>
<tr>
<th>Calcium antagonist</th>
<th>Vascular selectivity</th>
<th>Conduction system</th>
<th>Time to peak plasma levels</th>
<th>Elimination half-life</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nifedipine</td>
<td>++</td>
<td>-</td>
<td>20-40 min</td>
<td>2-4 h</td>
</tr>
<tr>
<td>Amlodipine</td>
<td>++++</td>
<td>-</td>
<td>6-12 h</td>
<td>35-50 h</td>
</tr>
<tr>
<td>Felodipine</td>
<td>++++</td>
<td>-</td>
<td>2-8 h</td>
<td>10-15 h</td>
</tr>
<tr>
<td>Isradipine</td>
<td>++++</td>
<td>-</td>
<td>1-2 h</td>
<td>7-8 h</td>
</tr>
<tr>
<td>Lacidipine</td>
<td>++++</td>
<td>-</td>
<td>3 h</td>
<td>7-18 h</td>
</tr>
<tr>
<td>Manidipine</td>
<td>++++</td>
<td>-</td>
<td>1-2 h</td>
<td>4-8 h</td>
</tr>
<tr>
<td>Nicalidipine</td>
<td>++++</td>
<td>-</td>
<td>1 h</td>
<td>4-5 h</td>
</tr>
<tr>
<td>Nimodipine</td>
<td>++++</td>
<td>-</td>
<td>2 h</td>
<td>7-18 h</td>
</tr>
<tr>
<td>Nimodipine*</td>
<td>++++</td>
<td>-</td>
<td>1-2 h</td>
<td>5 h</td>
</tr>
<tr>
<td>Nisoldipine*</td>
<td>++++</td>
<td>-</td>
<td>1-2 h</td>
<td>8-12 h</td>
</tr>
<tr>
<td>Nitrendipine</td>
<td>++++</td>
<td>-</td>
<td>2 h</td>
<td>8-14 h</td>
</tr>
<tr>
<td>Diltiazem</td>
<td>+</td>
<td>+</td>
<td>1-2 h</td>
<td>3-11 h</td>
</tr>
<tr>
<td>Verapamil</td>
<td>+</td>
<td>+</td>
<td>1-2 h</td>
<td>3-7 h</td>
</tr>
</tbody>
</table>

* Cerebral vessels > peripheral vessels
* Coronary vessels > peripheral vessels

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traction of human papillary muscles by 17 % whereas a concentration of 14 nMol already results in a 50 % contraction inhibition of isolated human coronary arteries [14]. These findings confirm amlodipine’s tissue selectivity in the human model as well. The guinea pig papillary muscle requires a five-fold higher amlodipine concentration compared to nifedipine to initiate a 50 % decrease in muscle contraction, which means that amlodipine has approximately a fivefold weaker negative inotropic action on the heart muscle compared to the parent compound nifedipine [15]. The relative tissue selectivity of amlodipine – mainly a consequence of different membrane potentials of heart and vascular muscles – is comparable to nitrendipine and is approximately fourfold to that of nifedipine [5]. The coronary dilating action of amlodipine and nifedipine is 3,000-fold stronger than papaverine compared to a factor of 10,000-fold for nisoldipine [15].

Binding studies of membrane preparations have shown that amlodipine binds and dissociates very slowly at the dihydropyridine receptor of the α1-subdivision of the calcium channel. Thus, binding saturation is not achieved until approximately five hours after administration compared to approximately one hour with isradipine [5]. A further special feature of amlodipine is that this substance interacts, to a certain extent, with the binding sites for phenylalkylamines and benzothiazepines. Plasma concentrations twenty- to thirty-fold above that required for vascular dilation are required to retard atrioventricular conduction, as demonstrated in animal models [16]. However, clinically relevant doses do not influence the contractility of the sinus node or atrioventricular conduction in humans [17].

The data of in vitro tests were confirmed in animal experiments. Amlodipine showed a gradual blood pressure (BP) lowering effect without reflex tachycardia. The maximum BP reduction was achieved approximately 8 hours post oral dosing and persisted for 24 hours, while the BP lowering effect with nitrendipine was no longer evident after 6 hours [13]. Long-term therapy over 30 weeks in SHR-rats blunted an increase in arterial BP and prevented the development of myocardial hypertrophy [5]. Cholesterol-enriched feeding tests in rats and rabbits have shown that additional administration of amlodipine reduces the development of atherosclerotic vascular lesions, with a dose-dependent effect [18]. The survival time of stroke–prone rats was prolonged [5]. Pretreatment with amlodipine also prevented the ischaemic calcium overload of myocytes in normotensive rats after reperfusion [5] and ischaemia-induced endothelial dysfunction [19]. Expansion of infarction zone and ventricular remodeling were not influenced [20]. However, Hoff et al. showed a decrease in the experimental infarction size in a group of canines treated with amlodipine [21]. There was no impairment of left ventricular function in dogs with acute myocardial infarction while on therapy with amlodipine [22]. Dogs pretreated with amlodipine and subjected to several ischaemic episodes with subsequent reperfusion experienced a more rapid recovery of regional wall movement, and a less-pronounced loss of energy rich phosphate in the ischaemic areas [23]. A study in pigs demonstrated beneficial effects in a model of pacing-induced heart failure with either amlodipine or fosinopril and combined amlodipine/angiotensin-converting enzyme inhibition (ACEI) provided greater benefit with respect to vascular resistance and neurohumoral activity compared with either monotherapy [24]. Overall, these findings from animal experiments highlight cardioprotective properties which would be desirable in the treatment of patients with coronary heart disease and/or arterial hypertension and probably heart failure.

Clinical pharmacology – therapeutic relevance

Following once daily oral administration, amlodipine is virtually completely absorbed from the intestinal tract. Peak plasma levels are reached after 6 to 12 hours. It has a relatively high bioavailability of 60 to 80 % [5, 25]. Steady state plasma concentrations in healthy volunteers are achieved after the 7th dose, without accumulation since this substance follows linear kinetics [13, 26]. Amlodipine has a very large volume of distribution (21 l/kg) and, similar to other CAs, 95 % are bound to plasma proteins. Amlodipine undergoes slow hepatic metabolism. Less than 10 % of the orally administered dose is excreted unchanged. The metabolites possess no calcium antagonistic properties and are excreted via urine (60 %) and in faeces (20–25 %). This drug has a very long elimination half-life of 35–50 hours [5, 25]. While patients with renal insufficiency showed no accumulation, elimination was delayed in patients with hepatic cirrhosis [27] so that monitoring of therapy is recommended in these patients. No adverse interactions were observed for digoxin, cinetidine, and nitroglycerine [5, 13]. Concurrent food intake does not influence amlodipine’s rate of absorption or plasma levels [28].

Studies over the last 10 years have shown that the incidence of myocardial infarction, sudden cardiac death, apoplectic insult and silent myocardial ischaemia show a circadian distribution of incidence with a peak in the early morning hours [29]. Quyyumi et al. have observed in patients with stable CHD using multiple ergometric exercise tests that the ejection fraction was lower in the early morning hours at the time of a significant ST segment depression than at mid-day or in the early evening [30]. Circadian rhythms were demonstrated for various physiological functions; heart rate and BP showed an increase in the early morning hours [31, 32]. Heightened sympathetic alpha-adrenoceptor activity associated with an increase in peripheral resistance was observed in normotensives [33] and hypertensives [34] in the early morning hours. Furthermore, clinical studies have shown that pharmacokinetics and pharmacodynamics of antihypertensives or antischismatic substances vary markedly with the time of the day, as observed with propranolol, organic nitrates and nifedipine [35]. In this complex situation, substances such as amlodipine are most beneficial because of a low variation of plasma levels, a high oral bioavailability, long elimination half-life and small variations between peak and trough plasma levels [26]. These properties insure a continuous efficacy over 24 hours with once daily dosing and hence improve patient compliance [36].

Clinical studies – coronary heart disease

The anti-anginal efficacy of varying amlodipine doses (1.25–10 mg) was investigated in 136 patients with stable angina pectoris compared to placebo. Exercise ECGs were carried out 24 hours after each dose [37]. Exercise duration was prolonged by 31 % with the 10 mg dose. Compared to baseline, a 48 % increase in the time to onset of angina was observed. The incidence of ischaemic attacks and use of glycerol trinitrate decreased significantly for all doses of amlodipine tested. In a study with similar design, once-daily administration of 10 mg amlodipine prolonged the time to onset of angina by 28 %; nitrate consumption decreased by 50 % and the frequency of anginal attacks dropped by 67 % [38]. Invasive measurements after a 20 mg intravenous application of amlodipine showed a reduction of systemic resistance with an increase in stroke volume and a slight increase in heart rate at rest and during
exercise. The maximum filling rate determined by radionuclide ventriculography was increased [39]. The maximum rate of pressure increase in the left ventricle remained unchanged after intravenous amlodipine administration (10–20 mg) during monotherapy and after pre-treatment with a β-blocker [17]. Bernink et al. [40] compared amlodipine and diltiazem in patients with stable exercise-induced angina in a double-blind study over a period of 8 weeks: 80 patients were assessed by means of ergometric tests 24 h after amlodipine and 12 h after diltiazem, respectively. The time to onset of angina was prolonged by 25% with amlodipine and by 3% with diltiazem compared to baseline. The decrease in the frequency of attacks and nitrate consumption were comparable in both groups. Silke et al. found a comparable reduction of angina while on antianginal treatment and a minimum of 4 hours and/or a total ischaemia time of ≥1 minute) documented by Holter monitoring over 48 hours and/or a total ischaemia time of ≥20 minutes [47]. In a 4-week study of 210 patients, BP was recorded both in the sitting and supine position at hourly intervals during the initial 12 hours and 24 hours post oral dosing. Diastolic BP was reduced in all patients at amlodipine doses ≥2.5 mg. Heart rate remained unchanged at all doses both in the lying and standing position compared to baseline values [49]. Parallel to amlodipine’s pharmacokinetic, the maximum BP lowering effect was observed after approx. 6–12 hours without any significant changes in heart rate. After discontinuation of therapy, baseline BP values were not attained even 6 days after cessation of therapy [25]. Ambulatory BP monitoring using intra-arterial [50] and non-invasive [51–54] measurements with observation periods of 4 and 28 weeks showed a continuous 24-h effect that BP control was present during sleep with preservation of circadian rhythm and no significant change in heart rate. The trough/peak ratio for the antihypertensive effect ranges from 50–100% with an average of 63% [55]. This represents a major advantage for long-term antihypertensive therapy in view of the aforementioned incidence of cardiovascular events in the early morning hours [29], a phase of the day during which short-acting antihypertensives fail to blunt rapid increases in BP upon waking [32, 34]. Hamada et al. evaluated 24-h ambulatory BP and simultaneous Holter monitoring before and after 4 weeks of administration of amlodipine, short-acting nifedipine, or its slow-release formulation [56]. While the mean hourly BP was reduced significantly and to similar degrees only nifedipine induced increases in heart rate, especially during the daytime. Invasive measurements have shown that the initial reduction of peripheral vascular resistance and the increase in stroke volume was unchanged one year after initiation of amlodipine therapy [52]. No loss of efficacy was observed in an open study of oral amlodipine over a period of 27 months [57].

Clinical studies – arterial hypertension

Over 200 patients with mild to moderate arterial hypertension were treated with amlodipine 1.25–10 mg in a placebo-controlled double-blind study over a period of 8 weeks [48]. The extent of BP reduction was dose-dependent and averaged -20/9 mmHg after 8 weeks with no changes in heart rate. In a similar 4-week study of 210 patients, BP was randomized and followed for 3 years. Compared to placebo amlodipine as add-on therapy reduced hospitalizations due to angina by 35% and revascularization procedures (PTCA or bypass) by 46%. For detailed analysis one has to await the final publication which hopefully will soon appear.

REVIEWS
for diastolic values, in patients with moderate to severe hypertension while on amiodipine. Amlodipine (5 mg/day) reduced systolic and diastolic BP to a similar degree as 8 or 16 mg/day of candesartancilexetil [65]. Enalapril and amloidipine were equally effective in patients with isolated systolic hypertension [66]. While short-acting DHGs should be abandoned due to serious adverse events long-acting DHGs are recommended by JNC VI report [67] as well as the 1999 WHO-ISH guidelines [68] as an alternative to diuretics for the first line therapy of isolated systolic hypertension.

Acebutolol (400 mg/day), amloidipine (5 mg/day), chlorothalidone (15 mg/day), doxazosin (2 mg/day) and enalapril (5 mg/day) were compared in the Treatment of Mild Hypertension Study (TOMHS). The final results showed similar efficacy in reducing systolic and diastolic blood pressure versus placebo for these antihypertensives in approximately 900 patients with mild hypertension [69]. Controversial findings were obtained with regard to concomitant therapy with diuretics. While Glasser et al. were able to achieve an additional BP reduction by adding on amloidipine to pre-existing diuretic therapy [70], bendroflumethiazide had no additional BP lowering effects compared to placebo after four-week pretreatment with amloidipine [71]. The administration of amloidipine in combination with captopril [72] or enalapril [73] resulted in an additional BP reduction of 18/12 mmHg and 19/10 mmHg compared to placebo. Coadministration of amloidipine and candesartan resulted in a significantly greater BP reduction than either drug alone [65].

Left ventricular hypertrophy (LVH) has been considered an independent risk factor for sudden cardiac death, ventricular arrhythmias, coronary heart disease and heart failure [74, 75]. The coronary reserve of hypertensive hearts is diminished by 30–50%. This implies a predisposition to myocardial ischaemia even in the haemodynamically-compensated state with normal coronary arteries [76]. Mortz et al. [77] were able to show an increase in the coronary reserve by approx. 30% in a small group of patients with angiographically normal coronary arteries who were treated over a period of 9–12 months with various antihypertensives. Furthermore, data from several studies indeed suggest a decrease in cardiovascular mortality and morbidity of hypertensive patients after LVIH regression [78–80]. In a more recent meta-analysis LVH which included predominantly clinical trials with short-acting DHGs had led to the conclusion that ACEI seemed to be more potent in this regard [81–83]. Recent publications which additionally included studies with long-acting CAs from 1990–1995 [84] and from mid 1995 to the end of 1996 [85] revealed that ACEI and CCBs were both superior to β-blockers and diuretics and equally effective in reducing LV muscle mass. The decrease in LV muscle mass was comparable in all treatment groups in the above-mentioned TOMHS study with the exception of chlorothalidone which showed the largest decrease in LV-mass, but also the largest decrease in LV end-diastolic diameter [69]. Several smaller studies have documented a decrease in LV muscle mass after treatment with 5–20 mg amloidipine for 3–12 months [86–88]. In one of our own studies, left ventricular muscular mass was reduced by approx. 9% following a 28-week amloidipine therapy together with an increase in early-diastolic filling [54]. This is comparable to earlier reports for chronic therapy with nitrépril or captopril [89]. Picca et al. showed a comparable decrease in LV muscle mass together with an increase in early-diastolic flow for both enalapril and amloidipine during the course of an 18-months observation period in patients with LVIH [90]. Regression of LV muscle mass was similar when amloidipine was compared to enalapril [91], fosinopril [92] or lisinopril [93] for 6–12 months. In addition, amloidipine was superior to lisinopril in previously untreated hypertensive patients in reducing the mean common carotid intima-media thickness [94] which is an established surrogate marker for early atherosclerosis.

Neurohormonal and metabolic effects

In vitro tests have shown that amloidipine reduces ADP- or collagen-induced thrombocyte aggregation [95]. The growth of human kidney mesangial cells is inhibited [96], as has been demonstrated for other calcium antagonists of the dihydropyridine group [5, 97]. The vasodilating effect of calcium antagonists in the kidneys predominantly affects afferent vessels. In experiments, the fall of glomerular filtration rate following administration of angiotensin II can be completely nullified with amloidipine [98]. In hypertensives, 4-week amloidipine therapy increased glomerular filtration rate by 11%, renal blood flow by 16% and decreased renal vascular resistance by 24% [99]. While Cappuccio et al. showed a stimulation of plasma renin activity in 6 patients with arterial hypertension and unchanged sodium excretion in 24-h urine during a 2-week treatment [100], Lund-Johansen et al. did not observe an increase in plasma volume or extracellular fluid in a larger group of amloidipine-treated patients [52]. Plasma renin activity, aldosterone and angiotensin II concentrations as well as plasma levels of atrial natriuretic peptide were unchanged in both young and elderly hypertensives [27, 54, 99–103].

In contrast to findings with the DHGs nifedipine [56], felodipine [104] or nitrendipine [105], amloidipine does not lead to a stimulation of noradrenaline and adrenaline secretion at rest or during exercise [27, 54, 56, 102, 103, 106, 107]. The absence of a neurohumoral stimulation is desirable especially during the pharmacotherapy of heart failure patients. In addition to the higher negative inotropic effect of older CAs, the recurrent stimulation of the sympathetic nervous system may explain the unfavourable patient outcomes in post-infarction trials with nifedipine [108]. A randomized, double-blind, placebo-controlled study of more than 100 patients with chronic heart failure in NYHA stages II–III and ejection fractions <40% has shown that the treatment with amloidipine in addition to a basis therapy with digoxin, diuretics and/or ACEI improved the exercise duration and heart failure symptoms within 8 weeks [106]. In the Prospective Randomized Amlodipine Survival Evaluation (PRAISE) more than 1100 patients with severe chronic heart failure and ejection fractions <30% were stratified according to ischaemic or nonischaemic aetiology and randomly assigned to double-blind treatment with either placebo or amloidipine while already on therapy with diuretics, ACEI and digitalis. While there was no difference in cardiovascular morbidity and mortality between the amloidipine and placebo groups among patients with ischaemic heart disease, amloidipine reduced the risk of death by 46% in patients with nonischaemic cardiomyopathy [109]. The latter finding needs confirmation which is being sought in the PRAISE-II trial. In this context, the cytoprotective activity of amloidipine which appears to be independent of its effects on calcium flux may play an important role by limiting the cytokine-induced damage in dilated cardiomyopathy [110]. In contrast to older DHGs which may have dangerous side effects in heart failure patients amloidipine is listed for the treatment of angina and hypertension in patients already on ACEI, diuretics, and digitalis [67].

After a 3-week amloidipine therapy (5 mg/day), no changes were monitored in insulin sensitivity, insulin secretion, total cholesterol, lipoprotein subfractions or triglycerides in the
The evaluation of approximately 40 placebo-controlled studies with roughly 3,000 patients showed a mild side effect profile typical for amlodipine’s peripheral vasodilating action (see Table 2). A placebo-controlled crossover study with a 2-week treatment period showed a significantly higher rate of side effects, in particular headache and flush, while on nifedipine in a retard preparation (41 %) versus amlodipine (27 %). With amlodipine, only the frequency of ankle edema was significantly higher (9 % vs. 2 %) when compared to placebo [60]. Side effects for once-daily doses of 20 mg nitratedipine were more frequent than for once-daily 5 mg amlodipine not only during the initial 3 days of therapy (39.5 % versus 5 %), but also at the end of the 4-week observation period (47.4 % versus 27.5 %) [61]. Compared to enalapril, both groups produced equally-effective BP reduction with a discontinuation rate of 4 % overall with cough (13 %) in the enalapril group and lower leg edema (22 %) in the amlodipine group as the most frequently reported side effects [63]. In a multicentre study of more than 100 patients, a side effect rate of 27.5 % was observed in out-patients over a period of 3 months with lower leg edema (13.8 %) as the most frequent adverse event [116]. Whereas in the CAPE study side effects occurred during amlodipine treatment in 17.3 % of the patients compared to 13.3 % for placebo [47], the side effect rate in the TIMHS study was higher in the placebo group than in any of the drug treatment group. Of all antihypertensives tested, amlodipine was best tolerated: 28.2 % of the initially randomized patients in the verum group still received this medication at the end of the 4-year observation period [69]. Clinical trial databases revealed incidence rates for all-cause mortality, myocardial infarction, and new/worsened angina among all amlodipine-treated patients of 3.0, 3.3, and 1.6/1000 patient-years of exposure, respectively [117]. Among those in comparative trials alone the all-cause death rate (3 out of 942 for a rate of 6.7/1000 patient-years) was comparable to that of non-CCB agents (4 out of 4126 for a rate of 4.1/1000 patient-years).

Last year two studies gained much interest in the context of the ongoing CCB controversy since they suggested that CAs probably promote adverse cardiovascular events in diabetic patients with hypertension. The Appropriate Blood Pressure Control in Diabetes (ABCD) trial [118] was a prospective, randomized, blinded comparison of the effect of moderately controlled BP (target diastolic BP 80–89 mmHg) with that of intensively controlled BP (target diastolic BP 75 mmHg) on the incidence and progression of complications in Type 2 diabetes using nisoldipine (10–60 mg/day) or enalapril (5–40 mg/day). After 67 months of the study the Data and Safety Monitoring Committee recommended the discontinuation of the hypertensive cohort since a significant difference in the rate of cardiovascular events was observed for patients on nisoldipine compared to enalapril. Data for the hypertensive subgroup (n = 470) on the incidence of myocardial infarction, a secondary endpoint of the study, were subsequently analyzed and published while the blinded treatment in the normotensive cohort (n = 480) was continued. Control of BP, blood glucose, and lipid concentrations were comparable in both treatment groups, but nisoldipine was associated with a higher incidence of fatal and nonfatal myocardial infarctions (25/235 vs. 5/235). Complications at base line were equally distributed between both treatment groups, however, there were significantly more patients on a concomitant β-blocker or diuretic therapy in the enalapril group.

The objective of the Fosinopril versus Amlodipine Cardiovascular Events Randomized Trial (FACET) was – somewhat contrary to what the title suggests – to compare the effects of open-label therapy with fosinopril (20 mg/day) and amlodipine (10 mg/day) on serum lipids and diabetes control in 380 Type 2 diabetics with hypertension which were followed for an average of 3.5 years [119]. While both treatments were equally effective in lowering BP with no differences in metabolic effects, the primary endpoint of the study, patients on amlodipine were at a significantly higher risk of acute myocardial infarction, hospitalized angina, and especially stroke. Of particular interest, however, is the fact that the risk of unfavourable cardiac events was lowest among patients (n = 108 or 28 % of the total study population) who were on combination therapy of fosinopril with amlodipine.

A comparison of the 5-year incidence rates for myocardial infarction in the aforementioned studies (amlodipine: 12 %, nisoldipine: 11 %, fosinopril: 9 %) to historical controls (Helsinki Heart Study: 7.5 % [120], Schwabing Study: 12.5 % [121]) or the results of the UK Prospective Diabetes Study Group 39 (captopril: 10 %, atenolol: 8 % [111]) suggests no real difference but rather a need to explain the unusually low incidence (2 %) in the enalapril treated patients in the ABCD trial. Also, subgroup analyses from the Systolic Hypertension in Europe (Syst-Eur) trial [123] revealed an even greater benefit for a DHP-based (nitratedipine) antihypertensive treatment in older diabetics with isolated systolic hypertension (all cardiovascular endpoints in diabetics -63 % compared to -21 % in non-diabetics). Finally, in the Hypertension Optimal Treatment (HOT) study [124] major cardiovascular events in the 1501 diabetic patients were halved with aggressive BP control (target diastolic BP ≤ 80 mmHg) which usually was achieved only by combination therapy with felodipine and ACEI and/or β-blocker compared to less intensive therapy (target diastolic BP ≤ 90 mmHg).

The PRAISE study [109] has established the safety of amlodipine for the treatment of angina and/or hypertension in patients with advanced left ventricular dysfunction. In the Antihypertensive and Lipid Lowering Treatment to Prevent Heart Attack Trial (ALLHAT) 40,000 patients have been randomized to compare amlodipine, lisinopril, doxazosin, and...
chlordiazepoxide for effects on nonfatal myocardial infarction or fatal coronary heart disease in high risk patients with hypertension. About 1/3 of the patients suffer from diabetes, however, after a separate evaluation of the primary endpoint in this subgroup representing more than 7000 patient-years the Data and Safety Monitoring Board of this study recommended that the trial continues according to the protocol [125].

In summary, once-daily amlodipine provides a favourable side-effects profile without adversely affecting neurohumoral or metabolic parameters, thereby providing safe and reliable 24-hour therapy for patients with coronary heart disease and/or arterial hypertension.

References


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