Cardiovascular Effects of NPK-1886, a New Dihydropyridine Compound with Calcium Entry Blocking Activity

Jun NAGURA, Bunsei MURAYAMA, Nobuo HARADA, Kunio SUZUKI, Tetsuji MIYANO, *Michio YAJIMA and *Kazumi TAKEYA

Central Research Laboratories, Banyu Pharmaceutical Co., Ltd., 2-9-3 Shimo-Meguro, Meguro-ku, Tokyo 153, Japan
*Department of Pharmacology, Aichi Medical University, Nagakute, Aichi 480-11, Japan

Accepted November 22, 1985

Abstract—The cardiovascular effect of NPK-1886 (NPK), a novel photostable dihydropyridine compound, was studied by comparing it with that of nifedipine (Nif). In normal Wistar rats (NWR), systolic blood pressure was only slightly depressed by NPK or Nif, while in three types of hypertensive rats (i.e., spontaneously hypertensive rats (SHR), renal hypertensive rats (RHR) and DOCA-saline-induced hypertensive rats (DOC-Na-R)), the hypotensive potency of NPK was more than or equal to that of Nif. The effectiveness of NPK on the normal and hypertensive models was in the following order: DOC-Na-R, RHR, SHR, NWR. Coronary perfusion flow in Langendorff’s heart was increased almost the same extent by NPK and Nif. On isolated rabbit aortic strips, the antagonistic potencies of NPK, like those of Nif, were greater for calcium than for norepinephrine, serotonin and angiotensin II. The negative ino- and chronotropic potency of NPK in isolated guinea-pig right atria was less than that of Nif. The slow membrane action potentials of guinea-pig papillary muscle were suppressed by NPK, but less than by Nif, with manifestations of a reduction of $V_{\text{max}}$ and AP-duration. These results indicate that NPK has a potent hypotensive effect on hypertensive models and a weaker cardiac inhibition. The general toxicity of NPK was lower than that of Nif.

Calcium entry blockers which block the calcium channels of cell membranes have significant effects on the cardiovascular system (1–9): they share a common ability to decrease pacemaking activity, reduce cardiac muscle force development, and relax vascular smooth muscle. In the heart, the calcium entry blockers have some crucial advantage for amelioration of the state of disordering of the circulation. They increase blood flows in the coronary vessel and other systemic vascular beds, diminishing afterload on cardiac muscle during blood ejection, and dispensing the oxygen consumption of the heart (10). In addition, calcium overload into the cell caused the death of the cell. Henceforth, calcium entry blockers were reported to protect the necrosis or mortal calcification of the cell (11, 12).

Nifedipine, a prototype calcium entry blocker was first reported by Bossert and Vater (13) and the mechanism of action was elucidated by Fleckenstein et al. (14, 15). Thereafter, many kinds of organic substances were introduced as a class of calcium entry blockers, but there was no common chemical structure essential for calcium entry blocking action (16). Because of considerable heterogeneity in the affinity of different cells for calcium entry blockers, some modification of the molecule of nifedipine yielded some different potencies on the blood vessel and heart, although the essential mechanism of action was the same (17).

An ideal calcium entry blocker should therefore have a low general toxicity and selective action on some specific part of the circulatory system and, in connection with
nifedipine, a photostable nature. With this aim, Miyano et al. (18) synthetized many dihydropyridine compounds, the number exceeding 300. Among them, we found NPK-1886 to be a compound ideal for antihypertension at the present stage. The chemical structure of NPK-1886 is isopropyl methyl 2-carbamoyloxymethyl-6-methyl-4-(2,3-dichlorophenyl)-1,4-dihydropyridine-3,5-dicarboxylate (Fig. 1). In this paper, we reported on the cardiovascular effects of NPK-1886, comparing them with those of nifedipine as a prototype dihydropyridine calcium channel blocking agent.

Materials and Methods

1. Measurement of blood pressure in rats: Normal Wistar rats (NWR) and hypertensive model rats (spontaneously hypertensive rats (SHR), renal hypertensive rats (RHR) and DOCA-saline-induced hypertensive rats (DOC-Na-R)) of the male sex, weighing from 250 to 350 g, were used. The animals were warmed-up in a box maintained at 37°C for 15 min prior to the experiments. The systolic pressure in the tail arteries of rats was measured in the conscious state using plethysmographical units connected with a tail-cuff (Ueda Co. VSM-10-S). Male SHR, 13–20 weeks of age, were used. RHR were prepared by the following methods: the left renal artery of rats (Wistar strain) under ether anesthesia was isolated and loosely ligatured by a silver wire at a slit width of 0.2 mm. Seven days after the operation, the residual right kidney was extirpated. Four weeks thereafter, the hypertensive state had developed, and the animals were then used. Rats of DOC-Na-R were prepared as follows: healthy male rats (Wistar strain) first received the left kidney extirpation at 5 weeks after birth. One week after the operation, 1% saline solution was administrated ad lib, and desoxycorticosterone acetate (DOCA) was injected hypodermally once a week in a dose of 5 mg/animal. The blood pressure monitoring was made 4 weeks after the first DOCA injection, and the rats in a hypertensive state of over 160 mmHg were used in the experiment.

2. Experiments on vascular system: To measure coronary flow, the heart was rapidly isolated from albino rabbits of either sex, weighing about 1.5 kg, and Langendorff’s heart was made by cannulating into the aorta. Well oxygenated Krebs-Henseleit solution (compositions in mM: NaCl, 115; KCl, 4.7; CaCl₂, 3.2; MgCl₂, 1.2; KH₂PO₄, 1.2; NaHCO₃, 24.9; and glucose, 10.0) at 30±1°C was perfused into the vascular bed of the heart. The negative logarithms of the concentration of test substances showing a 50% increase in perfusion rate was used as an index of the vasodilating potency.

Spirally cut strips (width 2 mm, length 20 mm) of blood vessel were made from the thoracic aorta of rabbits stunned by a blow on the head and exsanguinated, and the strips were suspended in a bath (37°C) at a resting tension of 1.5 g. The strips were first incubated for 1 hr in normal Tyrode solution (compositions in mM: NaCl, 137; KCl, 2.7;
CaCl₂, 1.8; MgCl₂, 1.1; Na₂HPO₄, 0.4; NaHCO₃, 12; and glucose, 5.6), and then the bath solution was replaced by Ca-free Tyrode solution and equilibrated for at least 30 min. Then the preparations were incubated in a Ca-free, K-rich Tyrode solution. Five min thereafter, CaCl₂ was added cumulatively into the solution. NPK-1886 or nifedipine was added 5 min before the experiment. Tension development in the strip was measured by means of a mechanoelectric transducer (Nihon Kohden SB-1T) which was connected to an amplifier and the data were recorded on oscillographic paper.

In experiments to determine the concentration-response curves of norepinephrine (NE), serotonin (5-HT) and angiotensin II (AT II) in the absence and presence of NPK-1886 or nifedipine, the development of tension in aortic strips was measured under the same conditions described above by adding these agents cumulatively into normal Tyrode solution in the bath.

3. Experiments on isolated cardiac muscle: Guinea pigs of either sex, weighing 250–300 g, were stunned by a sharp blow on the head and the hearts were rapidly excised. The right atria with pacemaker activity was isolated from the heart bathed in a well-oxygenated Krebs-Henseleit solution at 30°C. The preparation at a resting tension of 0.5 g was suspended in a bath containing the Krebs-Henseleit solution. The force of contraction was measured isometrically by means of a force-displacement transducer (Nihon Kohden, SB-1TH) and displayed on oscillographic paper after 1 hr equilibration in the bath. Heart rate was counted from the contraction frequency of the right atrial preparations.

Papillary muscles were isolated from the right ventricle of the excised heart and then fixed vertically or horizontally (in case of electrical activity measurements) in a bath containing normal Krebs-Henseleit solution of 30°C. The papillary muscles were stimulated by a rectangular pulse of 0.5 msec duration, at a strength of 1.3-fold threshold. The resting force was kept constant at 0.4 g. In normal cases, stimulation frequency was 1.0 Hz. In the case of partially depolarized papillary muscles, stimulation frequency was 0.1 Hz because of the severity of fully effective stimulation at a higher rate. The partially depolarized papillary muscle was made by increasing potassium concentration in Krebs-Henseleit solution up to 30 mM. Slow membrane action potentials elicited from the partially depolarized cells were recorded by conventional methods using the 3 M KCl-filled glass microelectrodes with a tip resistance of about 15 megohms. The parameters of slow action potential were analyzed with regard to the rate of rise ($\hat{V}_{\text{max}}$), amplitude (Amp) and duration (APD) (19, 20).

4. Acute toxicity: Lethal dose of NPK-1886 and nifedipine was determined by the Litchfield-Wilcoxon method from the data of experiments using healthy mice and rats. Mice (male and female) weighing 20±2 g and male rats weighing from 150 to 220 g were used. These two kinds of animals were fed on solid food (Crea, CA-1) and water ad lib. The animal room temperature and humidity were controlled at 22±2°C and 55±5%, respectively. No food except water was administered 24 hr before starting the experiments for oral administration of drugs.

5. Drug preparations: NPK-1886 and nifedipine were dissolved in polyethylene glycol 400 (PEG 400, Wako Pure Chem. Co.) and used in experiments evaluating the effects on blood pressure and the LD₅₀ in the i.v. route. In the cases of LD₅₀ estimation by administering the drugs in test animals in both i.p. and p.o. routes, NPK-1886 and nifedipine were suspended in 0.5% carboxymethyl cellulose (CMC) since they were not sufficiently soluble in PEG 400. In the in vitro studies, NPK-1886 and nifedipine were dissolved in dimethyl sulfoxide and then used. Experiments using nifedipine in vitro were performed in a dark room.

6. Statistical analysis: The values presented in the Figs. and Tables are expressed as the mean±S.E. For comparing two groups of results, statistical analyses were performed by Student’s t-test or the paired Student’s t-test.

7. Chemicals: The following drugs were used: nifedipine (Bayer), deoxycorticosterone acetate (DOCA, Wako Pure Chem. Co.), angiotensin II (Nakarai, Japan), norepinephrine (Daiichi Pharma. Co.), serotonin
creatinine sulfate (Nakarai, Japan), dimethyl sulfoxide (DMSO, Wako Pure Chem. Co.). NPK-1886 was synthetized as described in a previous report (18), and its purity was ascertained by elemental analysis, IR, UV, MS and NMR. NPK-1886 is a highly photo-stable substance. All the other chemicals used in these experiments were of special grade and provided from commercial sources.

Results

1. Effects on blood pressure: The time courses of the effect of NPK-1886 and nifedipine on the systolic blood pressure are shown in Fig. 2. In NWR, NPK-1886 in doses of 3–30 mg/kg, p.o., produced a mild lowering of blood pressure. The depressor effect of NPK-1886 was much the same as that of nifedipine.

In contrast, NPK-1886 produced a significant decrease in the blood pressure of SHR. Oral administration of NPK-1886 in doses of 3, 10, 30 mg/kg produced a significant decrease in systolic blood pressure dose-dependently. The maximum decrease was observed 1–3 hr after administration. Comparing the hypotensive potency of NPK-1886 and nifedipine, their dose-response curves at the maximum response during the observation (for 24 hr) were analyzed by the least squares method, and the dose of 30% decrease in blood pressure from the control level (ED30) were used as a measure of their potency. NPK-1886 was 1.4 times stronger than nifedipine; the ED30 values of NPK-1886 and nifedipine were 10.2 mg/kg and 14.3 mg/kg, respectively.

NPK-1886 and nifedipine significantly decreased the systolic blood pressure in DOC-Na-R of Wistar strain as well. The ED30 values of NPK-1886 and nifedipine in mg/kg were 0.7 and 0.8, respectively.

2. Effects on blood vessel: The effects of NPK-1886 and nifedipine on vascular perfusion through the heart were examined on Langendorff's rabbit heart. Test substances were injected into the cannula which was inserted to the aorta. The potency (as pD2 value) of NPK-1886 and nifedipine for increasing the flow rate by 50% was 7.09 ±0.06 (n=5) and 7.06±0.05 (n=20), respectively.

Concentration-response curves of aortic strips in a partially depolarized state by raising external K+ concentration to 40 mM are depicted for calcium concentration in the absence and presence of NPK-1886 and nifedipine. Both substances shifted the control curve to the right in a parallel fashion. The shift powers of NPK-1886 and nifedipine, evaluated from the pA2 value, were 9.64±0.06 (n=6) and 8.80±0.17 (n=5) (Fig. 3), respectively.

The antagonistic potencies of test substances for the other vasoactive agents, such as norepinephrine (NE), serotonin (5-HT) and angiotensin II (AT II), were examined as shown in Table 1. The maximum responses of vasoactive agents were taken as 100%. There were no appreciable differences in antagonistic potencies between NPK-1886 and nifedipine. The dose-response curve of NE shifted in a parallel fashion to the right in the presence of both substances like what was observed in the case of calcium ions. Both nifedipine and NPK-1886 depressed the contractile responses of aortic strips to 5-HT and AT II in a non-competitive manner (Fig. 4).

3. Effects on cardiac muscle preparations: Using the spontaneously beating atrial strips isolated from the guinea-pig heart, the chrono- and inotropic effects of NPK-1886 were examined, and the results were compared with those of nifedipine. Both NPK-1886 and nifedipine had negative ino- and chronotropic effects. Nifedipine was much more potent than NPK-1886 in both effects; NPK-1886 first caused 20% negative inotropy at a concentration level of 10^{-5} M, while nifedipine caused a similar effect at 10^{-7} M (Fig. 5, upper).

The potencies of the negative inotropic effect were also examined in the ventricular myocardium of guinea-pig papillary muscle driven at a frequency of 1 Hz. The negative
logarithms of the concentrations of NPK-1886 and nifedipine producing a 20% inhibition (pIC20) on the force of contraction were 6.08 and 6.97, respectively. Hence the potency of NPK-1886 was 1/8 that of nifedipine (Fig. 6).

With regard to the negative chronotropic effect, nifedipine was more potent than NPK-1886 (Fig. 5, bottom); the inhibition on the heart rate was first observed at $3 \times 10^{-6}$ M NPK-1886, which decreased the rate by 4.1%. 

**Fig. 2.** Time courses of the effects of NPK-1886 and nifedipine on systolic blood pressure after their oral administration in normotensive Wistar rats (NWR), spontaneously hypertensive rats (SHR), renal hypertensive rats (RHR) and DOCA-saline-induced hypertensive rats (DOC-Na-R). *, **: Significant difference from the control at $P<0.05$ and $P<0.01$, respectively.
while the rate decrease in the case of nifedipine at 3×10^{-7} M was 6.7%. Negative chronotropic potency of NPK-1886 was approximately 1/10 that of nifedipine.

Experiments were conducted to detect the inhibiting effects of NPK-1886 and nifedipine on the slow action potentials elicited from partially depolarized papillary muscles by increasing the external potassium concentration (30 mM). The potencies of these two compounds inhibiting the slow response were examined by conventional microelectrode methods. NPK-1886 inhibited the slow action potentials at a concentration of 10^{-5} M; at this concentration, NPK-1886 reduced significantly (by about 40%) the rate of rise (V_{max}), amplitude and duration of slow action potential. In contrast, nifedipine completely inhibited the slow action potentials at 10^{-7} M (Fig. 7).

4. Acute toxicity: The acute toxicity of NPK-1886 and nifedipine in mice and rats was examined and the results are summarized in Table 2. In mice, the general toxicity of NPK-1886 was low as evaluated from the LD50 values; in the intravenous route of NPK-1886 administration, the toxicity was 1/2 that of nifedipine. In the intraperitoneal route of administration, the toxicity of NPK-1886 was 1/4 that of nifedipine.

The LD50 values of both drugs orally administered to rats were more than 3000 mg/kg. In the intravenous route of administration, the general toxicity of NPK-1886 was less than 1/2 that of nifedipine. In addition, PEG 400, the vehicle of the drugs, did not have any toxic effect on mice or rats even when it was injected into the vein at a volume which was 4-fold that contained in the maximum dose administration.
Discussion

The present experiments clarified that NPK-1886 has a potent hypotensive effect on hypertensive animal models such as those with spontaneously developed, kidney-oriented and DOCA-saline-induced hypertensions. The hypotensive potency of NPK-1886 was estimated to be almost the same degree as that of nifedipine. Among the hypertension models used in the present experiments, DOCA-saline-induced hypertension was most effectively inhibited by both NPK-1886 and nifedipine. Renal hypertension was also highly effectively inhibited. In contrast, the blood pressure of normotensive rats was far less effectively reduced by both Ca-antagonistic agents. The reason why the compound was so selectively effective in hypertensive models in contrast to normotensive rats is now under discussion (21, 22). There is, therefore, a need to make further studies to clarify the mechanism by which these calcium channel inhibitors reduced more effectively the blood pressure of various hypertensive model rats than that of nor-
motensive ones.

In vascular smooth muscle cells, it is known that there are two mechanisms of calcium entry: one is receptor operated Ca-channels, and the other is voltage dependent Ca-channel (16). With regard to receptor operated Ca-channels, it is generally known that Ca-channel blockers such as nifedipine and verapamil are less effective for the contraction of vascular smooth muscle induced by vasoactive agents such as NE, 5-HT and AT II (16, 23, 24). This is the case in the present experiments using NPK-1886 and nifedipine as well.

In contrast to the receptor-operated mechanism, NPK-1886 and nifedipine exerted their action depending upon membrane potential; they were highly effective for both cardiac and vascular smooth muscles in the partially depolarized state. The contractile response of partially depolarized vascular smooth muscle to calcium ions was clearly inhibited by NPK-1886 and nifedipine. The present results can be interpreted as the indicating that NPK-1886 blocked voltage-dependent calcium channels in smooth muscle cell membrane.

It has been reported that among calcium entry blockers, there are tissue differences in the potency of calcium entry blocking action. For example, nifedipine was more potent for smooth muscle relaxation than for cardiac depression; In contrast to the concentration sufficient to relax the smooth muscle (3×10^{-9} M), nifedipine inhibits first the heart rate, contractility and impulse conduction at relatively higher concentrations in the order of 10^{-7} M, while verapamil acts on the heart and smooth muscle functions to the same extent at a given concentration (16, 25, 26).

In the present experiments, NPK-1886 blocked the slow action potentials of the myocardium first at relatively high concentration of 10^{-5} M, while nifedipine did that at 3×10^{-8} M. In view of the results, NPK-1886
Fig. 7. Inhibitory potencies of NPK-1886 and nifedipine on the slow membrane action potentials of papillary muscle isolated from the right ventricle of guinea pig. The muscle cell membrane was depolarized by raising the external KCl concentration to 30 mM. The maximum rate of rise ($V_{\text{max}}$), amplitude and duration (at a 90% repolarization level) of slow action potential are plotted against concentration of drug. Top figures, actual recordings of slow membrane action potentials recorded from a papillary muscle.

Table 2. Acute toxicities of NPK-1886 and nifedipine in mice and rats

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Sex</th>
<th>LD50 (mg/kg)</th>
<th>95% confidence limits</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>i.v.</td>
<td>i.p.</td>
</tr>
<tr>
<td>(MICE)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NPK-1886</td>
<td>male</td>
<td>38.0 (32.5–44.5)</td>
<td>3150 (2270–4380)</td>
</tr>
<tr>
<td></td>
<td>female</td>
<td>42.0 (36.5–48.3)</td>
<td>3300 (2540–4290)</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>male</td>
<td>17.0 (15.3–18.9)</td>
<td>970 (795–1180)</td>
</tr>
<tr>
<td></td>
<td>female</td>
<td>14.5 (12.7–16.5)</td>
<td>930 (762–1130)</td>
</tr>
<tr>
<td>(RATS)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NPK-1886</td>
<td>male</td>
<td>29.7 (26.4–33.4)</td>
<td></td>
</tr>
<tr>
<td>Nifedipine</td>
<td>male</td>
<td>13.0 (11.2–15.1)</td>
<td></td>
</tr>
</tbody>
</table>
can be estimated as being more selective for vascular smooth muscle than nifedipine is. This selectivity of NPK-1886 was also seen in the chronotropic effect (Fig. 5, bottom). The bathmotropic effect of NPK-1886 was not examined in the present study. However, NPK-1886 did not affect the electrocardiograph (ECG) of guinea pigs (H. Takeshita et al., unpublished data), suggesting that there is no appreciable effect on the conduction system.

Judging from the present studies, NPK-1886 is worthy of clinical trial for hypertensive disorders because of its selectivity and low general toxicity. In addition, because of the lack of a cardiac effect, NPK-1886 will improve cardiac failure by decreasing the afterload through the peripheral vasodilatatory effect.

Acknowledgement: We gratefully acknowledge the help given by Mr. H. Ando and other co-researchers in the Department of Pharmacology, Aichi Medical University.

References
12 Shen, A.C. and Jennings, R.B.: Kinetics of calcium accumulation in acute myocardial ischemic injury. Am. J. Pathol. 67, 441-452 (1972)
18 Miyano, T., Suzuki, K., Harada, N., Ushizima,


