Long-term effects with policosanol on lipid profile according to hypercholesterolemia severity in older patients

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Abstract

Background: End-point based studies have demonstrated a direct relationship between coronary disease and elevated serum levels of low density lipoprotein cholesterol (LDL-C) and total cholesterol, as well as the benefits of lowering LDL-C with statins on clinical end-points.

Policosanol is a mixture of very long chain fatty alcohols purified from sugar cane wax, with dislipidemia controlling effects, proved in numerous clinical assays in which patients with different conditions were included. The efficacy and tolerability of policosanol in the elderly have been also investigated in several clinical trials, being effective, safe and well tolerated.

Objectives: To investigate the effects of policosanol treatment during three years on lipid profile with a proportional intensity to the initial dislipidemia severity in older hypercholesterolemic patients.

Methods: The present analysis was obtained from the data of all patients treated with policosanol included in a previous prevention study. One thousand, four hundred seventy old patients of both sexes, between 60 to 85 years old, with type II hypercholesterolemia, and ≥ 1 non-lipid coronary risk factors, were randomized in two groups and treated with policosanol or placebo, during three years. Significant changes on lipid profile with a proportional intensity to the initial dislipidemia severity were considered primary efficacy variables. The analysis was done by Intention-to-treat method.

Results: An analysis of the response intensity show that after treatment, reductions of LDL-C, total cholesterol and triglycerides were greater, and according to the initial hypercholesterolemia severity, so that, patients with severe hypercholesterolemia showed the better responses, followed by moderate and mild hypercholesterolemia. An opposite pattern, however, was observed for HDL-C serum concentration. Triglycerides did not respond in the same way.

The frequency of vascular serious adverse events was lower in the policosanol group (15 events) compared with those on placebo group (49 events). There were 109 patients who experienced serious adverse events: 83 (11.3 %) in placebo and 26 (3.5 %) in policosanol group (p<0.0001). Twenty-three deaths occurred up to study completion: 19 patients belonging to placebo group (2.6 %) and 4 to the policosanol group (0.5 %).

Conclusions: The treatment with policosanol produce positive changes on serum lipid profile according to hypercholesterolemia severity and with a significant lower amount of vascular serious adverse events, mortality, and frequency of total adverse events in older patients.
1. Introduction

The management of cardiovascular risk factors, like elevated LDL-C, is called primary prevention when patients had not previously experienced a cardiovascular event. The focalization on primary prevention on LDL reduction is based on epidemiologic data documenting a continuous, positive and graded relationship between LDL-C concentration and cardiovascular disease events and mortality, evidencing that lowering LDL-C in patients, reduces the risk in patients with or without cardiovascular disease across a broad range of concentration [1-3].

Patients without known cardiovascular disease are generally at much lower baseline of cardiovascular events risk than patients with known cardiovascular disease. The decision to recommend LDL treatment depend on the global cardiovascular disease risk because the potential absolute risk reduction by treating hypercholesterolemia will usually be smaller for patients with established cardiovascular disease [2,3].

End-point based studies have demonstrated a direct relationship between coronary disease and elevated serum levels of LDL-C and total cholesterol[2], as well as the benefits of lowering LDL-C on clinical end-points [4-9].

Hypercholesterolemia management in the elderly had been questioned because, according with some opinions, elevated LDL-C and total cholesterol levels reduces its predictive value in ageing patients, because the relative decline of coronary risk with age [10]. However, plasma lipid determinations still remain as strong predictors for absolute coronary risk in the elderly [11], and the evidence obtained from strata analyses of older patients included in statin trials had shown the clinical benefits in this population [4-9].

On the other hand, increased HDL is considered a cardio protective factor. A large and prolonged multivariable analysis, had confirmed the inverse, independent, strong and graded relationship between HDL-C in both cardiovascular disease and coronary disease mortality [12].

Policosanol is a mixture of high molecular weight alcohols purified from sugar cane (Saccharum officinarum, L) wax [13], exhibiting a cholesterol-lowering effects due to the inhibition of cholesterol synthesis by regulating the activity of hydroxymethyl glutaryl Coenzyme (HMG CoA) through the increase of AMP kinase activity [14-17].

A total cholesterol, and LDL-C lowering, the HDL concentration increment and a constant but less effective reduction on triglycerides concentration of policosanol have been demonstrated in patients with type II hypercholesterolemia [18,19]. The efficacy and tolerability of policosanol in the elderly have been investigated in several clinical trials, being effective, safe and well tolerated in older individuals [20-27].

Policosanol shows many and relevant pleiotropic effects, such as the inhibition of platelet aggregation [28-30], and the susceptibility inhibition of LDL to be oxidised between many others [31,32]. Clinical studies and long-term post marketing surveillance studies have proven that policosanol is safe and well tolerated [13,18-35].

This background supported the conduction of a long-term study with policosanol in hypercholesterolemic elders with the objective to investigate its effects on lipid profile according to different degree of hypercholesterolemia severity in older patients during three years.

2. Patients and Methods

2.1. Study design

The data of hyperlipidaemia severity used in the present analysis on policosanol effect, was obtained from the results of all patients treated with policosanol included in a previous prospective, randomized, double-blinded, placebo-controlled study which included 1470 older patients [36], treated with placebo or policosanol for three years after randomization.

An Independent Ethics Committee approved the study protocol. Patients were recruited at four Polyclinic Centre and followed by a medical staff of the Surgical Medical Research Centre after providing informed written consent.
Patients were advised to follow a step one cholesterol-lowering diet for five weeks, after which lipid profile and safety laboratory indicators were assessed and the next week they attended to visit 2. Laboratory values obtained at the end of baseline period and safety physical indicators obtained at visit 2 were considered as baseline values for respective parameters.

2.1.1. Enrolment criteria

Patients of both sexes aged 60 to 80 with documented coronary disease, hypercholesterolemia, and others coronary risk factors were enrolled. The rationale for the lowest age was to include older subjects with an appropriate life expectancy which, in Cuba is around 80 years.

2.1.2. Inclusion criteria

Patients were included for randomization, if after the diet-only period, they showed total cholesterol $\geq 5.2$, LDL-C $\geq 3.4$ and triglycerides $<$4.52 mmol/L.

2.1.3. Exclusion criteria

Patients were excluded if active renal disease, diagnosed neoplastic disease, severe hypertension (diastolic blood pressure $\geq 120$ mm Hg), uncontrolled diabetes or poor cognitive function were present. In addition, patients who had had episodes of unstable angina, myocardial infarction, stroke or any serious adverse events within the three months previous to being enrolled in the study were also excluded.

2.1.4. Withdrawal criteria

Any serious adverse events or any adverse events justifying such decision, unwillingness to follow-up by any cause, major violations of study protocol, including $> 6$ consecutive weeks without taking the study medications.

2.2. Treatment

Tablets must be taken 5 mg once a day with evening meal. Patients should be titrated to 2 tablets oid if their total cholesterol levels after 6 or 12 months on therapy were $\geq 7$ mmol/L.

2.2.1. Compliance assessment

Compliance being assessed by patient questioning and tablet counts and defined as $\geq 85$ % of the scheduled tablets having been consumed since the prior visit.

2.2.2. Concomitant medications

Consumption of lipid-lowering drugs was forbidden from the time of enrolment to study completion, but no other restriction of concomitant therapy was done. Cases at secondary prevention were encouraged to take aspirin and/or $\beta$-blockers.

2.2.3. Assessments

Lipid profile and safety laboratory tests were performed at baseline and after 1, 2 and 3 years of randomization. At each visit dietary reinforcement and physical examination were done.

2.3. Efficacy analyses

2.3.1. Primary efficacy variables

Changes on lipid profile (LDL-C, total cholesterol, HDL-C and triglycerides) according to hypercholesterolemia severity were considered a primary efficacy variable in this analyses. Treatment was considered as effective if LDL-C was significantly reduced by $\geq 15$% [37].

2.3.2. Secondary efficacy variables

The incidence of vascular serious adverse events that occurred during the study was considered as a secondary efficacy variable. Vascular serious adverse events included all cardiovascular, cerebrovascular and peripheral events that led to the hospitalization or death of the patient.
2.3.3. Safety and tolerability analyses

Adverse event (AE) defined as any new unfavourable change in function, structure or laboratory data or the worsening of any pre-existing condition occurring through the study, independent of its relationship with treatment were considered in the safety and tolerability analysis.

AE were classified according to their intensity as mild, moderate or serious. Mild AE were those not requiring treatment or withdrawal of study medication, moderate AE required withdrawal of study medication and/or treatment of the AE [38].

Mild and moderate AE were also included for safety and tolerability analysis. Each AE was classified as having a causal relationship with treatment using the categories of definitely, probably, possibly, probably not, or definitively not drug-related.

Also, physical indicator (body weight, pulse rate, blood pressure) and laboratory test values (glucose, creatinine, aspartate aminotransferase –AST-, alanine aminotransferase –ALT-) were analysed.

2.3.4. Laboratory analysis

Blood samples were drawn after 12 hours overnight fasting at Policlinics and transported within the next 2 hours to the Surgical Medical Research Center for processing and analysis. Lipid profile and laboratory test values were determined by enzymatic methods using reagent kits (Roche). Laboratory analyses were performed in a Hitachi 719 autoanalyzer. Determinations were done on the same sampling day. A quality control was performed throughout the study, so that precision (within and between-day variations) and accuracy versus reference standards were controlled.

2.4. Statistical analysis

Statistical analysis for the whole study was planned in study protocol and amendments. All data were analysed according to Intention to treat principle, so that analyses were based on data of all randomised patients, as randomised.

Continuous values were compared using t test for paired (within group comparisons) and independent (between group comparisons) samples. Categorical data were compared with the $\chi^2$ test. All statistical tests were two-tailed, with significance at $\alpha=0.05$. Statistical analyses were performed using Statistics for Windows (Release 4.2; Copyright StatSoft, Inc. US) and SAS/STAT (Stat Soft, Version 8, US).

3. Results

3.1. Baseline patient characteristics

Of the 1612 patients recruited, 1470 were eligible and randomized to policosanol (n=737) or placebo (n=733). The main reasons to be not eligible were total cholesterol and LDL-C values after diet period below inclusion criteria (n=76); triglycerides > 4.52 mmol/L (n=36) and unwillingness to continue participating (n=30).

Table 1 summarizes the main baseline characteristics of study patients. Both groups were well matched at randomisation. Of 1470 randomised subjects, 466 (31.7 %) were at secondary prevention, while most were at primary prevention (1004, 68.3 %), but with ≥ 1 concomitant coronary risk factor. The prevalence of arterial hypertension, diabetes and current smoking was 64.1 %; 17.9 % and 20.2 %, respectively. Most patients (917, 62.3%) showed isolated hypercholesterolemia (elevated total cholesterol, normal triglycerides), while 553 (37.6 %) showed combined hypercholesterolemia (elevated total cholesterol and triglycerides).

Table 1 Main baseline characteristics of study patients

| Characteristics            | Placebo (n = 733) | Policosanol (n=737) |
|----------------------------|------------------|---------------------|
| Age (years) (X±SD)         | 66 ± 6           | 66 ± 6              |
| Body mass index (kg/m²) (X±SD) | 26.5 ± 5.3       | 26.7 ± 4.8          |
| Total cholesterol (mmol/L) (X±SD) | 6.70 ± 0.87      | 6.76 ± 0.90         |
| LDL-C (mmol/L) (X±SD)      | 4.65 ± 0.86      | 4.72 ± 0.88         |
| HDL-C (mmol/L) (X±SD) | 1.21 ± 0.33 | 1.22 ± 0.34 |
|----------------------|-------------|-------------|
| Triglycerides (mmol/L) (X±SD) | 2.23 ± 0.99 | 2.23 ± 0.90 |
| Systolic blood pressure (mm Hg) (X±SD) | 136 ± 17 | 137 ± 20 |
| Diastolic blood pressure (mm Hg) (X±SD) | 82 ± 10 | 82 ± 10 |
| Gender: Female | 582 | 79.4 | 571 | 77.5 |
| Male | 151 | 20.6 | 166 | 22.5 |
| Isolated hypercholesterolemia | 465 | 63.4 | 452 | 61.3 |
| Combined hypercholesterolemia | 268 | 36.6 | 285 | 38.7 |

### Risk factors

| Arterial hypertension | 473 | 64.5 | 470 | 63.8 |
| Smoking | 152 | 20.7 | 145 | 19.7 |
| Coronary disease* | 201 | 27.4 | 207 | 28.1 |
| Diabetes mellitus | 132 | 18.0 | 131 | 17.8 |
| Obesity (kg/m² > 30) | 66 | 9.0 | 63 | 8.5 |
| HDLc< 0.9 mmol/L | 51 | 7.0 | 59 | 8.0 |
| Cerebrovascular disease** | 34 | 4.6 | 36 | 4.9 |
| Family history of coronary disease | 362 | 49.4 | 366 | 50.0 |

### Concomitant medications (CM)***

| Diuretics | 181 | 24.7 | 187 | 25.4 |
| Calcium antagonists | 158 | 21.6 | 155 | 21.0 |
| Aspirin | 129 | 17.6 | 119 | 16.1 |
| Anxyolytics | 118 | 16.1 | 121 | 16.4 |
| β-blockers | 107 | 14.6 | 98 | 13.3 |
| Vasodilators | 95 | 13.0 | 90 | 12.2 |
| Oral hypoglycemic drugs | 79 | 10.8 | 65 | 8.8 |

n Number of patients; X mean, SD standard deviation, *myocardial infarction, unstable angina, coronary surgery. **stroke, ischemic transient attacks; ***CM consumed by > 6% of study patients.

### 3.2. All comparisons were not significant

Table 2 lists the frequency of withdrawals, which was greater (p<0.0001) in placebo group (189, 25.8 %) than in policosanol group (88, 11.9 %). The same was true (p<0.0001) for withdrawals due to adverse events and other reasons, these last ones being mainly related with patients showing alert values (total cholesterol ≥ 9.0 mmol/L). Two hundred and seventy-seven patients (18.8 %) withdrew from the study. Of them, 109 discontinued because of serious adverse events and another 12 (9 placebos, 1.2 % and 3 policosanol treated, 0.4 %) because of mild or moderate adverse events.

Compliance within the study drugs was good, since 721/737 (97.8 %) policosanol patients and 715/733 (97.5 %) of placebo adhered to compliance criterion (> 85 % of dose taken at the end of treatment) during the time that they received treatment.

Most policosanol patients (665/737, 90.2 %) were treated with 5 mg/d during the study. Three hundred eight (308) patients: 72 (9.8 %) policosanol and 236 placebos (32.2 %) were titrated to 2 tablets oid with the evening meal. The frequency of needing titration was different in both groups (p<0.01).
Table 2 Withdrawal analysis

| Withdrawals due to AE                          | Placebo (n = 733) | Policosanol (n = 737) | p value* | Total |
|------------------------------------------------|-------------------|-----------------------|----------|-------|
| Vascular SAE                                   | 49                | 15                    | p < 0.0001 | 64    |
| SAE from other causes                          | 34                | 11                    | p < 0.001 | 45    |
| Mild and moderate AE                           | 9                 | 3                     | ns       | 12    |
| Subtotal due to all AE                         | 92                | 29                    | p < 0.0001 | 121  |

Withdrawals due to other reasons

| Reasons                                                      | Placebo (n = 733) | Policosanol (n = 737) | p value* | Total |
|--------------------------------------------------------------|-------------------|-----------------------|----------|-------|
| Unsatisfactory efficacy                                      | 16                | 13                    | ns       | 29    |
| Travels abroad+changes to other towns or living areas       | 30                | 30                    | ns       | 60    |
| Unwillingness to follow-up                                  | 14                | 9                     | ns       | 23    |
| Subtotal due to other reasons                               | 97 (13.2 %)       | 59 (8.0 %)            | p < 0.01 | 156 (10.6 %) |
| Total of withdrawals                                        | 189 (25.8 %)      | 88 (11.9 %)           | p < 0.0001 | 277 (18.8 %) |

3.3. Effects on primary efficacy variables

Table 3 shows the effects of policosanol on lipid profile variables according to the severity of hypercholesterolemia at baseline. These data revealed a difference from placebo response of LDL-C, total cholesterol, HDL-C and triglycerides in all strata of hypercholesterolemia severity.

Table 3 Long-term effects of policosanol on lipid profile (x ± SD) according to hypercholesterolemia severity in older patients

| Study groups               | Baseline | 1 year | 2 years | 3 years |
|----------------------------|----------|--------|---------|---------|
| **Total cholesterol (TC) (mmol/L)** |          |        |         |         |
| **Mild hypercholesterolemia (TC ≥ 5, but < 6.1)** |          |        |         |         |
| Policosanol (n=200)        | 5.79 ± 0.22 | 5.21 ± 0.53* | 5.19 ± 0.64* | 5.16 ± 0.67* |
| Placebo (n=207)            | 5.76 ± 0.22 | 5.97 ± 0.55 | 5.99 ± 0.50 | 5.93 ± 0.52 |
| **Moderate hypercholesterolemia (TC ≥ 6.1, but < 7.8)** |          |        |         |         |
| Policosanol (n=424)        | 6.80 ± 0.47 | 5.71 ± 0.59* | 5.41 ± 0.60* | 5.26 ± 0.57* |
| Placebo (n=435)            | 6.80 ± 0.48 | 6.64 ± 0.63 | 6.72 ± 0.63 | 6.72 ± 0.60 |
| **Severe hypercholesterolemia (TC ≥ 7.8)** |          |        |         |         |
| Policosanol (n=113)        | 8.31 ± 0.55 | 6.25 ± 0.75* | 5.78 ± 0.89* | 5.42 ± 0.66* |
| Placebo (n=91)             | 8.33 ± 0.38 | 7.80 ± 0.91 | 7.88 ± 0.84 | 7.82 ± 0.84 |
| **LDL-C (mmol/L)**         |          |        |         |         |
| **Mild hypercholesterolemia (TC ≥ 5, but < 6.1)** |          |        |         |         |
| Policosanol (n=200)        | 3.89 ± 0.45 | 3.33 ± 0.49* | 3.16 ± 0.63* | 3.13 ± 0.59* |
| Placebo (n=207)            | 3.85 ± 0.45 | 4.11 ± 0.58 | 4.18 ± 0.55 | 4.14 ± 0.54 |
| **Moderate hypercholesterolemia (TC ≥ 6.1, but < 7.8)** |          |        |         |         |
| Policosanol (n=424)        | 4.77 ± 0.59 | 3.77 ± 0.59* | 3.42 ± 0.59* | 3.19 ± 0.58* |
| Placebo (n=435)            | 4.75 ± 0.59 | 4.71 ± 0.67 | 4.86 ± 0.65 | 4.85 ± 0.65 |
### 3.4. Intensity of response versus degree of dislipidemia

An analysis of the responses intensity of the showed that reductions in LDL-C, total cholesterol and triglycerides after one year of treatment were greater according to the degree of hypercholesterolemia severity, so that patients with severe hypercholesterolemia showed the better responses, followed by moderate and mild hypercholesterolemia. An opposite pattern, however, was observed for HDL-C. With the exception of triglycerides, the other responses were improved during the study. These changes are summarized and simplified in Tables 4 and 5 showing absolute values and percent changes, respectively.

### 3.5. Effects on secondary efficacy variables

The frequency of vascular serious adverse events was lower in the policosanol group (15 events) as compared with placebo (49 events) (p<0.0001) (Table 6).

The amount of cardiovascular serious adverse events compared to placebo (33) was significantly lower in the policosanol group (7) (p<0.0001). Also, there were 12 cerebrovascular serious adverse events (1.6 %) in the placebo and 5 (0.7 %) in the policosanol group (p<0.05).

There were 109 patients who experienced serious adverse events (fatal + non-fatal): 83 (11.3 %) in placebo and 26 (3.5 %) in policosanol group (p<0.0001).
Twenty-three deaths occurred up to study completion: 19 taking placebo (2.6 %), and 4 policosanol (0.5 %). The frequency of deaths due to cardiovascular events with policosanol (1 death, 0.1 %) was lower (p < 0.01) than with placebo (13 deaths, 1.8 %). Also, 3 patient taking placebo (0.4 %), but any patient under policosanol treatment died because of cerebrovascular events. The deaths due to nonvascular causes (6/23, 26.1 %) were similar in both groups.

On the other hand, the frequency of non-vascular serious adverse events in the policosanol group (11 events, 1.5%) was significantly lower (p <0.001) than in placebo (34 events, 4.6%) (data not shown in Table for simplicity).

### 3.5.1. Safety and tolerability

Policosanol did not modify safety indicators (data not shown in Table for simplicity). Thus, it did not raised ALT, AST, glucose or creatinine values, body weight and pulse rate were unchanged, but systolic and diastolic pressure were significantly (p<0.0001) reduced compared with baseline and placebo.

**Table 4** Summary of the changes on lipid profile variables in policosanol group according to hypercholesterolemia (HC) severity

| Study groups | Baseline | 1 year | 2 years | 3 years |
|--------------|----------|--------|---------|---------|
| **Total cholesterol (mmol/L) (x ± SD)** | | | | |
| Mild HC (n=200) | 5.79 ± 0.22 | 5.21 ± 0.53 | 5.19 ± 0.64 | 5.14 ± 0.67 |
| Moderate HC (n=424) | 6.80 ± 0.47 | 5.71 ± 0.59 | 5.41 ± 0.60 | 5.26 ± 0.57 |
| Severe HC (n=113) | 8.31 ± 0.55 | 6.25 ± 0.75 | 5.78 ± 0.89 | 5.42 ± 0.66 |
| **LDL-C (mmol/L)** | | | | |
| Mild HC (n=200) | 3.89 ± 0.45 | 3.33 ± 0.49 | 3.16 ± 0.63 | 3.13 ± 0.59 |
| Moderate HC (n=424) | 4.77 ± 0.59 | 3.77 ± 0.59 | 3.42 ± 0.59 | 3.19 ± 0.58 |
| Severe HC (n=113) | 5.98 ± 0.77 | 4.23 ± 0.71 | 3.71 ± 0.88 | 3.21 ± 0.62 |
| **HDL-C (mmol/L)** | | | | |
| Mild HC (n=200) | 1.15 ± 0.31 | 1.28 ± 0.23 | 1.34 ± 0.29 | 1.35 ± 0.23 |
| Moderate HC (n=424) | 1.21 ± 0.33 | 1.27 ± 0.24 | 1.31 ± 0.24 | 1.40 ± 0.21 |
| Severe HC (n=113) | 1.35 ± 0.39 | 1.36 ± 0.28 | 1.39 ± 0.28 | 1.48 ± 0.28 |
| **Triglycerides (mmol/L)** | | | | |
| Mild HC (n=200) | 2.00 ± 0.81 | 1.67 ± 0.54 | 1.72 ± 0.38 | 1.74 ± 0.53 |
| Moderate HC (n=424) | 2.24 ± 0.90 | 1.80 ± 0.57 | 1.82 ± 0.50 | 1.77 ± 0.30 |
| Severe HC (n=113) | 2.61 ± 0.95 | 1.88 ± 0.60 | 1.84 ± 0.43 | 1.90 ± 0.62 |

On the other hand, the report during the study, of mild and moderate adverse events was also significantly lower in the policosanol group than in the placebo group (p <0.01) (data not shown in Table for simplicity).

**Table 5** Lipid profile % changes in policosanol group (according to hypercholesterolemia- HC- severity)

| Study groups | 1 year | 2 years | 3 years |
|--------------|--------|---------|--------|
| **Total cholesterol** | | | |
| Mild HC (n=200) | -11 | -11 | -11 |
| Moderate HC (n=424) | -16 | -20 | -23 |
| Severe HC (n=113) | -25 | -30 | -35 |
| **LDL-C** | | | |
| Mild HC (n=200) | -14 | -19 | -20 |
| Moderate HC (n=424) | -21 | -28 | -33 |
|                  | Placebo (n = 733) | Policosanol (n = 737) | p value* |
|------------------|-------------------|-----------------------|----------|
| Primary efficacy | n                 | %                     | n        |
| Vascular SAE     | 49                | 6.7                   | 15       | 2.0     | p < 0.0001 |
| Cardiovascular SAE| 33                | 4.5                   | 7        | 0.9     | p < 0.0001 |
| Cerebrovascular SAE| 12              | 1.6                   | 5        | 0.7     | p < 0.05   |
| Secondary efficacy |                  |                       |          |
| SAE (fatal + non-fatal) | 83            | 11.3                  | 26       | 3.5     | p < 0.0001 |
| Non vascular SAE | 34                | 4.6                   | 11       | 1.5     | p < 0.01   |
| All mortality    | 19                | 2.6                   | 4        | 0.5     | p < 0.001  |

SAE serious adverse events, *Comparison with placebo (χ² test)

### 4. Discussion

The present analysis demonstrates that long-term treatment with policosanol produce positive changes on lipid profile according to hypercholesterolemia severity in older patients.

Both groups were well balanced at baseline. Most subjects were at primary prevention with one or more risk factors, but secondary prevention patients with a generally stable condition were also included. Hence, the study results should be extrapolated mainly to patients with similar conditions.

Among the most relevant baseline characteristics must be noted that the mean age of study patients was 66 years at randomization, indicating that many subjects still were young enough to apply preventive measures that might improve their quality and expectancy of life. The frequency of concomitant medications was high, consistent with their risk condition and common in the elderly [11].

LDL-C is considered the most important variable among lipid profile parameters. As compared with placebo, policosanol reduced LDL-C, total cholesterol and triglycerides, whereas it increased HDL-C. These changes were consistent with those expect response to policosanol, being potentially useful for risk reduction [12,17-26].

The responses were maintained, or even enhanced, throughout the study. The changes here reported for LDL-C, total cholesterol and HDL-C are consistent with the expected response to policosanol long-term therapy. Reductions on triglycerides, however, were superior that those reported in previous studies, a finding without any conclusive explanation. No significant change of any lipid profile variable occurred in placebo group.
The analysis of the cholesterol-lowering response according to hypercholesterolemia severity at randomization revealed that reductions in LDL-C, total cholesterol and triglycerides after one year of treatment were greater in severe and moderate hypercholesterolemia compared with mild hypercholesterolemia, while the opposite pattern was observed for HDL-C. These findings, agree with previous results. With the exception of triglycerides, which showed a similar response throughout the trial, the other responses were enhanced during the study when we analyze the data according to hypercholesterolemia strata.

The different withdrawal rate in both groups was a consequence of the discontinuations due to serious adverse events and those due to unsatisfactory efficacy for achieving levels over those considered as upper cut-off for premature discontinuations. Thus, the frequency of all vascular serious adverse events, cardiovascular, cerebrovascular, all deaths to vascular causes and all deaths was lower (p<0.05) than in placebo, consistently with LDL-C lowering and pleiotropic effects of policosanol, all beneficial for vascular function, thus preventing the occurrence of vascular events.

It was demonstrated that policosanol inhibits cholesterol synthesis in the first step of its metabolic pathway through activation of Adenosine Monophosphate Protein Kinase (AMPK), which in turn inhibit Hydroxyl-Methyl-Glutaryl-Coenzyme A-Reductase [13-16]. AMPK, once activated, also inhibit Acetyl CoA Carboxylase (ACC). The inhibition of ACC increases fatty acid oxidation and reduces lipid synthesis, protecting in this way, muscle, heart, and others tissues from lipotoxicity [39]. In addition, AMPK activation is associated with a wide array of beneficial effects, [40]. that could explain the low level of side effect and compliance in the treated group versus placebo.

After intestinal absorption, very long chain fatty alcohols are up taken by the liver and partially converted into carboxylic acids [41]. These results indicated that higher intake of VLCFA is significantly associated with favorable metabolic status including lower levels of circulating triglycerides [42]. Other study confirmed that circulating serum VLCSFAs were independently associated with favorable profiles of blood lipids (lower triglycerides and increase HDL-C); others cardiovascular disease risk markers, and a lower cardiovascular disease risk by 52 % [43].

On the other hand, fatty alcohols are substrates for the synthesis of plasmalogens in peroxisomes, which are potent endogenous antioxidants. Plasmalogens are released from the liver as component of lipoproteins thus protecting them from oxidation, and favoring its functionality [44].

Thus, the contribution of other effects, beyond its lipid-lowering properties, must be present in the benefits here demonstrated for policosanol. In particular, the contribution of its antiplatelet effects could be relevant, taking into account the effects reported for antiplatelet therapy on risk reduction in patients at high vascular risk [45]. Moreover, according with recent results, policosanol seems to present regeneration abilities via enhancement of HDL functionality [46].

Policosanol was safe and well tolerated. Unexpectedly, policosanol reduced nonvascular serious adverse events, a finding that could to be explained in basis of its pleiotropic effects, including those described and others yet unknown. The analysis of the overall frequency of any adverse event also discards any increase in particular adverse events due to policosanol.

Overall frequency of mild and moderate adverse event was lower in the policosanol group than in the placebo group. This result, together with serious adverse events and withdrawal analysis, eliminates any increase in particular adverse event due to policosanol.

No drug-related impairment of any safety indicator was observed. Policosanol, not placebo, modestly, but significantly reduced blood pressure, consistently with some previous data.22,24,25 Such decreases could have contributed to the presents results, since lowering systolic pressure significantly reduces coronary events and total mortality in the elderly. [47]. Specific on the effect of policosanol on hypertension studies confirm these results [48-50].

5. Conclusion

Elder patients with documented coronary disease, hypercholesterolemia, and others coronary risk factors, once treated with policosanol, reported relevant positive changes on serum lipid profile and a significant lower amount of vascular serious adverse events, frequency of total adverse events and mortality.
Compliance with ethical standards

Financial Disclosure
This study was support by the National Centre for Scientific Research, as part of its research-development projects.

Conflict of interest
The authors declare that there is no conflict of interest in relation to this work.

Statement of informed consent
Informed consents was obtained from all individual participants included in the study.

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