Sulodexide therapy for the treatment of diabetic nephropathy, a meta-analysis and literature review

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Abstract: Sulodexide is a heterogeneous group of sulfated glycosaminoglycans (GAGs) that is mainly composed of low-molecular-weight heparin. Clinical studies have demonstrated that sulodexide is capable of reducing urinary albumin excretion rates in patients with type 1 and type 2 diabetes, suggesting that sulodexide has renal protection. However, this efficacy remains inconclusive. In this article, we used meta-analysis to summarize the clinical results of all prospective clinical studies in order to determine the clinical efficacy and safety of sulodexide in diabetic patients with nephropathy. Overall, sulodexide therapy was associated with a significant reduction in urinary protein excretion. In the sulodexide group, 220 (17.7%) achieved at least a 50% decrease in albumin excretion rate compared with only 141 (11.5%) in the placebo. The odds ratio comparing proportions of patients with therapeutic success between the sulodexide and placebo groups was 3.28 (95% confidence interval, 1.34–8.06; P=0.01). These data suggest a renoprotective benefit of sulodexide in patients with diabetes and micro- and macroalbuminuria, which will provide important information for clinical use of this drug as a potential modality for diabetic nephropathy, specifically, the prevention of end-stage renal disease that is often caused by diabetes.

Keywords: sulodexide, diabetic nephropathy, meta-analysis, odds ratio

Introduction

The increased prevalence of diabetes has also led to an increase in the number of macro- and microvascular complications of diabetes, such as coronary heart disease, stroke, visual impairment, diabetic kidney disease, and end-stage renal disease. Diabetic nephropathy (DN) is a multifactorial complication with long-term consequences of chronic renal insufficiency.1 It is one of the major causes of end-stage renal disease and is associated with increased cardiovascular morbidity and mortality.2 Diabetic kidney disease was previously known as DN and is defined as diabetes with albuminuria (ratio of urine albumin to creatinine ≥30 mg/g), impaired glomerular filtration rate (<60 mL/min/1.73 m²), or both, and is the single strongest predictor of mortality in patients with diabetes.3 However, diabetic kidney disease encompasses not only DN but also atheroembolic disease, ischemic nephropathy, and interstitial fibrosis that occurs as a direct result of diabetes.4 Therefore, early identification of intensive therapy for DN is urgently needed. In addition to oxidative stress and hemodynamic changes, glycosaminoglycans (GAGs) are another factor contributing to the onset of glomerular abnormalities in diabetic patients.1 The thickening of glomerular basement membrane (GBM) and the depletion of GAGs cause decreased electrostatic charge barrier in the pathophysiology of DN.3 A novel therapeutic approach for DN is the reestablishment of heparin sulfate synthesis by GAG drugs.6 GAG replacement therapy for DN began 20 years ago, and sulodexide is the most extensively studied of the GAGs used to reduce albuminuria in diabetic patients.7,8 Sulodexide is composed of low-molecular-weight...
heparin and dermatan sulphate. Glycosaminoglycan sulodexide may affect the morphology and function of the basement membranes in microvessels. Sulodexide may protect against DN initiation and progression, as manifested by the reduced albuminuria. However, two collaborative trials have adequately assessed the efficacy of sulodexide in patients with type 2 diabetes mellitus (DM) and failed to demonstrate renoprotection for both micro-(incipient) and macro-(overt) albuminuria. Therefore, we performed this meta-analysis to investigate the effect of sulodexide on the progression of proteinuria in DN patients.

**Material and methods**

**Search strategy**

We searched PubMed, MEDLINE, and Web of Science, Scopus, and EMBASE in April 2015, using the following search terms: “diabetic nephropathy”, “proteinuria”, “albuminuria”, and “sulodexide”. We screened the publications by titles first, then the abstracts. We then evaluated the full-text version for inclusion and exclusion criteria, after exclusion of nonrelevant publications and identifications of duplicates from the different databases. The languages of publication were restricted to English and Chinese. All clinical studies except case reports were chosen. All searched data were retrieved. Authors’ bibliographies and references of selected studies were also searched for other relevant studies.

**Selection criteria**

In this meta-analysis, we collected all eligible articles about the relationship between sulodexide and micro-/macroalbuminuria and clinical outcomes in DN. The following inclusion criteria were applied: (1) type of study design was a randomized controlled trial that compared sulodexide with placebo, no treatment, or other antihypertensive drugs (excluding angiotensin-converting enzyme inhibitors [ACEIs] and angiotensin receptor blockers [ARBs]) on the effect of urinary albumin excretion; (2) all the participants were diabetic patients with proteinuria; (3) the study reported the changes in urinary protein excretion from baseline; and (4) study duration was longer than 3 weeks. The following exclusion criteria were applied: (1) articles that had no information on proteinuria or that could not be calculated by the albumin excretion ratio of means from the given information; (2) case reports, letters, reviews, expert opinion, conference abstracts, editorials, and non-English- and non-Chinese-language papers; and (3) all articles using cell lines, human xenografts, and in vitro/ex vivo studies.

**Data extraction**

The eligible studies were extracted by two investigators independently. Disagreements were resolved by discussions and consensus. The following information was recorded for each study: the first author name, year of publication, number of cases, sample source, micro- and macroproteinuria, and clinicopathological parameters. Two investigators reviewed all the articles that fit the inclusion and exclusion criteria. Heterogeneity of data was evaluated to determine whether or not the data of the various studies could be analyzed. Data for study characteristics and clinical response were summarized and represented in table format.

**Statistical analysis**

We used RevMan 5.2 (The Nordic Cochrane Centre, Copenhagen, Denmark) and the Stata 12.0 (Stata Corporation, College Station, TX, USA) for this analysis. Comparisons of dichotomous measures were determined by pooled estimates of odds ratios (ORs) as well as their 95% confidence intervals (CIs). Heterogeneity was determined by a chi-square test, with significance being set at $P<0.10$; the total variation among studies was estimated by $I^2$, with significance being set at $I^2>50\%$. A $P$-value of $<0.05$ was considered to be statistically significant. We used a random-effect model to pool the OR when there was heterogeneity among studies; otherwise, a fixed-effect model was selected. Studies reported albuminuria or proteinuria in different units that could not easily be compared. We referred to the methods by Kunz et al and we chose to summarize the therapeutic effects of sulodexide on urinary protein excretion using the ratio of the average therapeutic effects in the intervention group relative to the control group. This is roughly comparable across different measurement units and can be directly applied in a clinical context.

We first collected 64 articles from PubMed, MEDLINE, and the Web of Science, Scopus, and EMBASE. Finally, 13 full-text studies were retracted for more detailed assessment, after initial screening of the titles and abstracts for eligibility. Of these, three crossover trials without placebo group as control were excluded. Eventually, several publications were selected and met the inclusion criteria for this meta-analysis. The article search process and study selection are showed in Figure 1.

**Results**

**Study characteristics and quality**

Ten eligible studies published from 2005 to 2013 were finally selected for this study. A total of 2,770 patients,
including type 1 and type 2 diabetic patients from People’s Republic of China, Australia/New Zealand, the Netherlands, Poland, Spain, United Kingdom, Canada, and the United States, were enrolled. Eventually, several publications were selected and met the inclusion criteria for this meta-analysis. Their basic characteristics and antiproteinuric effects are summarized in Table 1. The confounding factors in the placebo groups and sulodexide groups, such as patient number, age, baseline proteinuria parameters, and mean arterial pressure before treatment, are summarized in Table 2. It was shown that the confounding factors that influence urinary albumin excretion were well-balanced in the sulodexide and control groups. All these possible confounders were well-controlled in the cited works and did not affect the meta-analysis.

Antiproteinuric effects of sulodexide (urine albumin excretion)

Although the different primary diseases (type 1 or type 2 diabetes), stages of proteinuria (micro- or macro albuminuria), follow-up periods, and races may affect the effects of sulodexide on proteinuria, we combined all the data for an analysis due to limited information under each specific situation. We took the maximum maintenance phase of follow up (not including the washout phase) in each study if multiple follow-up periods were available and compared the mean ratios of urine albumin excretion rate (AER) between the sulodexide and placebo groups (baseline proteinuria levels before therapy vs those after therapy). Overall, sulodexide therapy was associated with a significant reduction in urinary protein excretion. The ratio of means comparing the means of patients between the sulodexide and placebo groups from eight studies was: 0.76 (95% CI, 0.62–0.93, \( z = 2.63, P = 0.009 \)) (Figure 2). We conducted a random-effect meta-analysis, since \( I^2 = 70\% \). We removed one study at a time to analyze sensitivity and found the pooled ratio of means was not significantly changed, indicating the quality of the studies was acceptable.

Then, we compared the proportions of patients achieving the therapeutic success, defined as at least a 50% decrease in AER when treated with sulodexide. This criterion was similar to those used in the study by Heerspink et al, which...
Table 1  Clinical trials testing antiproteinuric effects of sulodexide in diabetes patients

<table>
<thead>
<tr>
<th>Patients (n)</th>
<th>Mean age*, y</th>
<th>Type of patients</th>
<th>Baseline proteinuria*</th>
<th>Mean arterial pressure before treatment, mmHg</th>
<th>Dose</th>
<th>Duration of treatment</th>
<th>Main results</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>54</td>
<td>32.1</td>
<td>Type 1 DM</td>
<td>24.5 g/d</td>
<td>&lt;120–123.9</td>
<td>600 lipoprotein lipase-releasing units/day IM</td>
<td>3 weeks</td>
<td>Significant reduction of albuminuria in 90% of patients, slower in macroalbuminuric patients</td>
<td>Dedov et al</td>
</tr>
<tr>
<td>223</td>
<td>47.4</td>
<td>Type 2 and type 1 DM</td>
<td>38.9 g/d</td>
<td>136.1</td>
<td>50 mg/d, 100 mg/d, or 200 mg/d PO</td>
<td>4 months</td>
<td>Dose-dependent reduction in albumin excretion rate</td>
<td>Gambaro et al</td>
</tr>
<tr>
<td>43</td>
<td>34.6</td>
<td>Type 1 DM</td>
<td>126.1 mg/d</td>
<td>N/A</td>
<td>120 mg/d PO</td>
<td>6 months</td>
<td>Reduction in albuminuria and NAG excretion, increase in renal vascular function</td>
<td>Sulikowska et al</td>
</tr>
<tr>
<td>149</td>
<td>64.1</td>
<td>Obese type 2 DM</td>
<td>124.8 mg/d</td>
<td>130</td>
<td>200 or 400 mg/d PO in addition to ACEI or ARBs</td>
<td>6 months</td>
<td>23.3% and 33.3% of the patients respectively in the two sulodexide groups combined and in the 200 mg/d group achieved a significant reduction or normalization of albuminuria vs 15.4% of the patients in the control group (P=0.06 and P=0.07, respectively)</td>
<td>Heerspink et al</td>
</tr>
<tr>
<td>62</td>
<td>60.3</td>
<td>Type 2 DM</td>
<td>N/A</td>
<td>134.4</td>
<td>600 lipoprotein lipase-releasing units/day IV</td>
<td>3 weeks</td>
<td>Significant decrease in urinary albumin excretion rate</td>
<td>Kang</td>
</tr>
<tr>
<td>986</td>
<td>59.0</td>
<td>Type 2 DM</td>
<td>2.9 g/d</td>
<td>98–102</td>
<td>200 mg/d, PO</td>
<td>250 lipoprotein lipase-releasing units/day, PO</td>
<td>Any change in albuminuria was identical in sulodexide group and control group</td>
<td>Lewis et al</td>
</tr>
<tr>
<td>1,248</td>
<td>62.0</td>
<td>Type 2 DM</td>
<td>N/A</td>
<td>130.7</td>
<td>200 mg/d, PO</td>
<td>200 mg/d, PO</td>
<td>Did not detect any significant differences between sulodexide and placebo</td>
<td>Packham et al</td>
</tr>
<tr>
<td>75</td>
<td>45.5</td>
<td>Type 2 DM</td>
<td>2.03 g/d</td>
<td>N/A</td>
<td>600 lsU IV</td>
<td>20 days</td>
<td>Albuminuria was reduced</td>
<td>Bu et al</td>
</tr>
<tr>
<td>55</td>
<td>62.8</td>
<td>Type 2 DM</td>
<td>2.23 g/d</td>
<td>N/A</td>
<td>600 lsU IV</td>
<td>3 weeks</td>
<td>24-hour urine protein, FIB, and β2-MG were decreased</td>
<td>Liu</td>
</tr>
<tr>
<td>65</td>
<td>50.0</td>
<td>Type 2 DM</td>
<td>600.2 mg/d</td>
<td>140</td>
<td>600 lsU IV</td>
<td>2 weeks</td>
<td>Relieved albuminuria effectively</td>
<td>Xiong</td>
</tr>
</tbody>
</table>

Note: *Data are expressed as the mean value, unless indicated otherwise.

Abbreviations: β2-MG, β2-microglobulin; ACEI, excluding angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; DM, diabetes mellitus; FIB, fibrinogen; IM, intramuscular; IV, intravenous; LSU, livestock unit; N/A, not available; NAG, N-acetyl-beta-D-glucosaminidase; PO, per os (orally); y, years; d, day.
Table 2 The confounders selected and matched from ten included studies

<table>
<thead>
<tr>
<th>References</th>
<th>Patients (n)</th>
<th>Age* (y)</th>
<th>Initial status of the kidney</th>
<th>Mean arterial pressure before treatment mmHg</th>
<th>Difference in mean arterial pressure mmHg</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Packham et al</td>
<td>62.8</td>
<td>No</td>
<td>2.03</td>
<td>122.7</td>
<td>2.59</td>
<td>*</td>
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<tr>
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<td>62.8</td>
<td>No</td>
<td>2.03</td>
<td>122.7</td>
<td>2.59</td>
<td>*</td>
</tr>
</tbody>
</table>

Notes: Data are expressed as the mean value, unless indicated otherwise. AER, albumin excretion rate; SCR, serum creatinine; ACR, albumin to creatinine ratio; N/A, not available; CRF, chronic kidney disease; OR, odds ratio; RR, relative risk. The ratio of means, comparing the means of patients between the two groups was 1.03 (95% CI, 0.89–1.19, $P=0.69$) (Figure 4).

Changes in serum creatinine and creatinine clearance

There were no significant differences in the change of serum creatinine and creatinine clearance between the two groups. The ratio of means, comparing the means of patients between the sulodexide and placebo groups was 1.03 (95% CI, 0.89–1.19, $P=0.69$) (Figure 4).

No heterogeneity was observed in the analysis of the effects of sulodexide on AER (Figure 2) and serum creatinine clearance (Figure 4), so the fixed-effect model was used. There was a heterogeneous found in the analysis of the effects of sulodexide on the achievement of therapeutic success (Figure 3), so the random effect model was used.
Side effects

Four trials briefly described the adverse events.\textsuperscript{11,12,18,21} The incidence of “likely related” side effects was similar in the sulodexide (ranging from 10.9\% to 35.2\%) and placebo groups (ranging from 12.2\% to 32.3\%). No serious adverse events were believed to be related to the study medication. The most common complaints were cardiovascular disorders; renal and urinary disorders; gastrointestinal and hepatobiliary disorders; infections and infestations; respiratory, thoracic, and mediastinal disorders; injury; poisoning, and procedural complications; metabolism and nutrition disorders; neoplasms (benign, malignant, and unspecified); musculoskeletal and connective tissue disorders; general disorders; administration-site conditions, eye disorders; nervous system disorders; vascular disorders; endocrine, reproductive, and breast disorders; blood and lymphatic system disorders; and ear and labyrinth disorders.\textsuperscript{11,12,18,21}

Publication bias and sensitivity analyses

A sensitivity analysis was conducted to assess the result stability, using the method in which one study was removed at a time. The pooled ratio of means and ORs were not significantly changed, indicating the stability of our study. The funnel plots were largely symmetric (Figure 5), suggesting...
there were no publication biases in the meta-analysis of sulodexide and clinicopathological features.

**Discussion**

DN occurs in 20% to 40% of patients with DM and is the major cause of morbidity and mortality in patients with type 1 and type 2 DM. The earliest clinical evidence of nephropathy is an increase in microalbuminuria, defined as more than 30 mg/day into the macroalbuminuria defined as more than 300 mg/day. As researchers have gained a clearer understanding of the pathophysiological mechanisms of DN, several new therapeutic agents have been subjected to clinical trials, including anti-TGF-β monoclonal antibody, anticonnective tissue growth factor monoclonal antibody.

As a component of the GBM, the heparin sulfate proteoglycans (GAGs) strongly influence the permeability of the glomerular filtration barrier by affecting its thickness and...
A decrease in the GAG composition of the GBM, particularly in heparin sulfate, is associated with diabetic kidney diseases. The loss of GBM integrity could worsen proteinuria and accelerate the progression of end-stage renal disease. Sulodexide is a GAG extract of the porcine lung and liver, and its major components are low-molecular-weight heparin and dermatan sulfate. Many preliminary data demonstrated that orally administered sulodexide was able to reduce urine albumin excretion in diabetic patients. However, in two large randomized double-blinded placebo-controlled trials led by Lewis et al and Packham et al which involved around 1,000 subjects with type 2 DM with micro- and macroalbuminuria in each trial, sulodexide failed to reduce urine albumin excretion compared with placebo. Therefore, the use of sulodexide for diabetic therapy was considered as “another one bites the dust”, to be abandoned.

Gambaro and Coccheri, who also led clinical trials of the effects of sulodexide on DN, disagreed with the conclusions of Lewis et al. They argued that discounting the efficacy of sulodexide in DN was premature. They interpreted the negative results obtained by Lewis et al differently: First, the drug used in the Collaborative Study Group trial by Lewis et al was only tested for heparin fraction activity without consideration for the dermatan sulfate, which constitutes one-fifth of the active ingredients of the drug. Second, treatment with maximal doses of concomitant drugs, such as ACEIs and ARBs, left little allowance for a superimposed effect of sulodexide. Third, patients in the Collaborative Study Group trials had more severe DN than did those in previous trials. In this meta-analysis, we showed that sulodexide therapy was associated with a significant reduction in urinary protein excretion (Figures 2 and 3) – both in reduced AER and proportion of subjects achieving therapeutic success.

Sulodexide has a renoprotective effect through restoration of glomerular ionic permselectivity, but the exact mechanism is still unclear. Several mechanisms have been proposed, one of which is dependent on GBM permselectivity, where sulodexide reduces proteinuria and improves renal function via inhibition of PKC-βIII, ERK, FGF-2, and heparanase-1, preventing epithelial–mesenchymal transition. Another proposed mechanism is independent of GBM permselectivity, in which sulodexide improves endothelial dysfunction through reduction of vascular endothelial growth factor.

In addition, a recent experimental animal model study has indicated that sulodexide may protect early but not late nephropathy (radiation and DN). Taken together, it is too early to say that sulodexide is ineffective in the treatment of diabetic nephropathy – the findings from both clinical trials and experimental models should be viewed as justifiﬁcation for continuing to develop this drug into clinical application.

Disclosure
The authors report no conﬂicts of interest in this work.

References


