Sodium-Glucose Co-Transporter 2 Inhibitors could Improve the Bioavailability of Vitamin C at the Kidney in Diabetes Treatment

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Letter to the Editor

It is known that plasma (serum) vitamin C level is lowered in diabetic patients by some mechanisms including renal loss [1]. Vitamin C and glucose filtered from the glomerulus are presumed to be reabsorbed at the renal proximal tubule by sodium-vitamin C co-transporter (SVCT) 1 and sodium-glucose co-transporter (SGLT) 2, respectively [2-4]. In the enterocyte that expresses SVCT 1 and SGLT 1, it was demonstrated in vitro that cellular uptake of vitamin C was inhibited by increasing concentrations of glucose in medium, which was not observed at the presence of phlorizin, a non-specific SGLT inhibitor [2]. If such an effect is applied to renal proximal tubular cells, it is hypothesized that SGLT 2 inhibitors can improve the bioavailability of vitamin C in diabetes treatment.

In two cases of diabetic patients treated with empagliflozin, a SGLT 2 inhibitor, in hospitalization, their serum vitamin C levels (normal reference range, 5.5 μg/ml to 16.8 μg/ml) have been observed to increase soon after its administration. In a male patient newly diagnosed with type 2 diabetes (HbA1c, 13.4 %) at age 41 (174 cm, 114 kg), his serum vitamin C level changed from 7.2 μg/ml to 8.1 μg/ml two days after adding 10 mg empagliflozin on 1000 mg metformin. Urinary glucose excretion increased from 15.5 g/day to 148.1 g/day. In a female patient with 12-year duration of type 2 diabetes (HbA1c, 7.4%) and renal insufficiency (serum creatinine, 1.64 mg/dl) at age 67 (155 cm, 99 kg), her serum vitamin C level changed from 7.9 μg/ml to 8.5 μg/ml twelve days after adding 10 mg empagliflozin on 5 mg linaglitin. Urinary glucose excretion increased from 2.9 g/day to 23.8 g/day. Ascorbic acid, a reduced form of vitamin C, or glucose enters cells specifically through SVCT or SGLT, driven by the inward sodium gradient maintained with sodium pump (Na⁺/K⁺-ATPase). Dehydroascorbic acid, an oxidized form of vitamin C, enters or exits cells competitively with glucose through facilitative glucose transporters (GLUTs) [3,5]. Taken together, a model of putative cellular transport of vitamin C in the early proximal tubular cells is drawn in Figure 1.

Ascorbic acid reabsorbed at the brush border membrane through SVCT 1 is oxidized by reactive oxygen species and exits the cell as dehydroascorbic acid competitively with glucose through facilitative glucose transporter (GLUT) 2 in the basolateral membrane. SVCT 1 and sodium-glucose co-transporter (SGLT) 2 are driven by the inward sodium gradient maintained by sodium pump (Na⁺/K⁺-ATPase).

Figure 1 A model of putative cellular transport of vitamin C in the early proximal tubular cell and its molecular structures. Vitamin C reabsorbed as ascorbic acid in the brush border membrane through vitamin C transporter (SVCT) 1 is oxidized by reactive oxygen species and exits the cell as dehydroascorbic acid competitively with glucose through facilitative glucose transporter (GLUT) 2 in the basolateral membrane. SVCT 1 and sodium-glucose co-transporter (SGLT) 2 are driven by the inward sodium gradient maintained by sodium pump (Na⁺/K⁺-ATPase).
GLUT 2 by becoming less competitive with glucose. Under the reducing and neutral pH conditions in the bloodstream, almost all vitamin C exists as ascorbic acid (precisely as ascorbate) [6]. It seems plausible that reabsorbed ascorbic acid protects renal proximal tubular cells from reactive oxygen species and exits the cells as dehydroascorbic acid, as indicated in the neurons [7]. Most recently, it has been demonstrated that SGLT 2 inhibitors slow progression of kidney disease in type 2 diabetes [8,9], where improvement of the bioavailability of vitamin C is not included in the putative mechanisms behind their renal effects. If this hypothesis is verified, SGLT2 inhibitors would be expected to have beneficial effects on the kidney still unknown beyond glucose lowering.

Conflict of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

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