DAILY TREATMENT WITH SILDENAFIL REVERSES ENDOTHELIAL DYSFUNCTION AND OXIDATIVE STRESS IN AN ANIMAL MODEL OF METABOLIC SYNDROME

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OBJECTIVES
Fructose consumption might be a contributing factor to the development of obesity and the accompanying cardiovascular disorders (hypertension, ...) seen in the metabolic syndrome

Baron et al., Am J Cardiol (1999); Anderson et al., J Am Coll Cardiol (1995)

Patients with metabolic syndrome exhibit generalized endothelial dysfunction

DeSouza et al., Diabetes Care (2002); Rosano et al., Eur Urol. (2005)

Moreover, since oxidative stress has been suggested to contribute to insulin resistance and associated endothelial dysfunction, we sought to determine the effects of chronic sildenafil on a potentially relevant biomarker of endothelial dysfunction, urinary 8-isoprostanes (IPT) content, a direct marker of non-enzymatic in vivo lipid peroxidation and the most reliable and clinically relevant marker of oxidative stress available to date.

MATERIALS & METHODS
Experimental animals
Wistar rats (n=10-14 per group) were fed a standard chow (CONT) or a 60% fructose/5% fat (% by weight)-enriched diet for 8 weeks (FFR). From week 5 through 8, sildenafil was administered twice a day (sc, 20 mg/kg, FFR+SIL), thus reaching clinically relevant plasma concentrations circa 20 nM unbound known to give efficacy in man (Pfizer Inc., data on file), then a 1-week wash-out period from sildenafil was observed.

In vitro evaluation of endothelial function
Isometric tension studies were performed on isolated aortic and superior mesenteric arterial (SMA) rings precontracted with phenylephrine to build concentration-response curves (CRC) to endothelium-dependent (ACh and A23 187) and -independent (SNP) relaxants in presence of indomethacin. Vascular cGMP content, urinary excretion of nitrates and nitrates (NOx) and 8-isoprostanes (IPT), and plasma levels of IL-6 and TNF-α were also evaluated.

RESULTS

No influence of the fructose-enriched diet on the animal body weight, blood glucose levels, or basal mean arterial pressure (MAP)

AUC of %increase of Relaxation (% Phe)

Urinary IPT (pg/ml/24h)

Urinary NOx excretion levels after one-week of wash-out from chronic sildenafil treatment

Enhanced compensatory endothelium-independent relaxations to SNP in FFR were not modified by sildenafil treatment

Significantly reduced endothelial relaxation responses to Ach (and A23187) in FFR in presence of indomethacin in both aortas and SMA

Restoration of endothelial relaxations induced by Ach (and A23187) in FFR treated chronically with sildenafil in both aortas and SMA

Neither the fructose diet nor the sildenafil treatment modified significantly :
- Tissue basal cGMP and ET-1 content in homogenates of aortas and SMA
- Urinary NOx excretion levels after one-week of wash-out from chronic sildenafil treatment
- IL-6 and TNF-α plasma levels

CONCLUSIONS

Effect of chronic sildenafil treatment on physiological parameters relevant to the metabolic syndrome : Correction of the enhanced response to glucose overload, as well as the hypertriglyceridemia induced by fructose feeding

Effect of chronic sildenafil treatment on in vitro vascular reactivity : Restoration of normal endothelium-dependent relaxations to Ach and A23 187 in aortas and mesenteric arteries

Effect of chronic sildenafil treatment on urinary 8-isoprostanes excretion : Normalization of the biological marker for oxidative stress

The beneficial effects of daily sildenafil on vascular endothelial dysfunction gives additional insight to the possible mechanism of action of sildenafil in cardiovascular disorders related to the metabolic syndrome that could be explored in future clinical trials. Moreover, this study provides preclinical support for the predictive value of endothelial reactivity and associated biological marker of oxidative stress such as urinary IPT as surrogate markers in future clinical trials addressing cardiovascular risks.