Bladder and erectile dysfunctions in the type 2 Diabetic Goto-Kakizaki rat

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OBJECTIVES

- >Urological functional complications such as bladder and erectile dysfunctions (ED) significantly impact the quality of life of diabetic
- Most of experimental in vivo studies of ED/bladder dysfunction caused by diabetes have used type 1 diabetes models
- A robust model for type 2 diabetes urological complications is lacking.

Aim of the study:

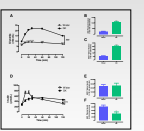
- Evaluate bladder and erectile function in the Goto-Kakizaki (GK) rat model for type 2 diabetes
- Evaluate the responses to standard-of-care treatments for overactibe bladder and erectile dysfunction in GK

general features of the animals

	Wistar	GK
Metabolic parameters		
Body weight (g)	415.1±5.8	356.8±4.1***
Glycemia (mmol/L)	6.13±0.17	9.71 ± 0.63***
Insulinemia (pmol/L)	264.12±25.20	299.40 ± 46.15
Triglycerides (mmol/L)	2.05±0.14	1.46 ± 0.09**
HDL cholesterol (mmol/L)	1.25±0.03	1.74 ± 0.03***
Total cholesterol (mmol/L)	1.66±0.05	2.14±0.04***
FFA (mmol/L)	0.55±0.02	0.47 ± 0.03
Diuresis (ml/24h)	10.6± 0.6	10.4 ± 0.9
Testosterone (ng/mL)	5.15±0.85	3.10±0.25°

MDL: high density lipoproteins; FFA: free fatty acids.

Data are the mean ± SEM of n= 25 Wistar rats and n= 23 GK rats. "p< 0.05,
""p<0.01, """p< 0.001, versus age-matched Wistar rate. Student's t. b. ...



Blood glucose (A) and blood insulin levels (D) in 18 weeks GK rats and in age-matched Wister rats 0, 10, 20, 30, 60 and 120 min after oral glucose challenge (Eg/s) to day weight). Area under the curve (AUC) of plucose or insulin time course curves from to 12 100 minutes (B and E, respectively) or 10 s0 minutes (C and F, respectively). Data are mean s SEM of experiments performed in Wister rats (n = 2 minutes) and K oft and (n = 20 a minute). \$4870.001 time-way ANOVA with

expected, diabetic GK rats showed hyperglycemia hyperinsulemia, hypercholesterolemia, and impaired glucose tolerance. In accordance with previous studies describing this model GK rats used in the current study represent a suitable model of type 2 diabetes to investigate urological complications.



Summary of research design

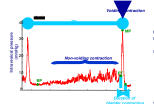
xperimental groups					
Groups	Strain	Cystometry	Erectile function experiments		
		acute iv treatment	acute iv treatment		
Control	Wistar	Vehicle (n=12) or	Vehicle (n=9) or		
		Solifenacin (n=9)	sildenafil (n=8)		
Diabetic	Goto-Kakizaki	Vehicle (n=9) or	Vehicle (n=10) or		
	(10 mode)	Soliferacio (n10)	eldenafil (n=11)		

•Male GK rats (n=25, GK/Par colony) and age-matched Wistar rats (n=23) were used between 14 and 19 weeks of age depending on the function evaluated. Metabolic parameters of these non obese GK rat model of type 2 diabetes compared to Wistar rats have been characterized at 14-week-old

MATERIALS & METHODS

Cystometry experiments
Cystometric investigation was performed in conscious rats. A bladder catheter, implanted 48h before experiment, was connected to a pressure transducer for bladder pressure monitoring and to a syringe-pump for bladder perfusion. The bladder was continuously perfused (50 µl/min) with saline. After stabilization, solifenacin (1 mg/kg) or its vehicle was administered by i.v route (250-300 µL during 1 min) and intravesical pressure was recorded during another 60 min period. The following parameters were analyzed: michurition pressure (mmHn)

duration (s) and AUC (mmHg x s) of voiding contraction; basal pressur (mmHg); pressure threshold at which voiding is initiated (mmHg); intercontraction interval (s); bladder capacity (μl, infusion rate x intercontraction interval), voided volume (μl) and voiding efficiency (%, as the ratio of voided volume/infused volume x100). The amplitude (mmHq) and the frequency (contraction per minute) of the non-voiding bladder contractions during the filling phase with an amplitude of >3 mm Hg were analyzed as well as the volume threshold to elicit NVC (percent of total



Frectile function evaluation: electrical stimulation of the cavernous nerve

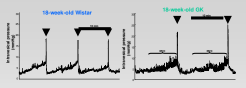
After 5 minutes of baseline recording of simultaneous computerized measure of mean arterial pressure (MAP) and intracavernous pressure (ICP), the CN was stimulated (6 V, 1 ms for 45 s) at different frequencies (0, 2,5,5,7,5,10, 12,5 and 15Hz) at 3-minute intervals in a randomized manner in order to assess the erectile responses. Erectile responses to ES CN were expressed as a ratio of ICP (mmHg) / MAP (mmHg) x 100, ICP being the difference between ICP in the flaccid state, i.e. before stimulation and ICP during the plateau phase of the erectile response, and MAP, the mean arterial pressure during the plateau phase, and as the ratio of AUCtot / MAP with AUCtot, with the area under the curve during the entire erectile response, measured from the beginning of the electrical stimulation until the end of the erectile response and determined using the ICP level in the flaccid state before the onset of the stimulation

Statistical analysis

All results were presented as mean ± SEM. Statistical analysis for general features, metabolic and urodynamic parameters were performed using Student's t-test. For OGTT and erectile function evaluation, comparisons of frequency-response curves were performed with a two-way ANOVA statistical analysis test followed by a Bonferroni's post-test. In case of significant interaction between the two factors (i.e. frequency of ES CN and experimental group), the difference between groups of rats will be examined by the modified Student's t-test with the Bonferroni's adjustment for multiple comparisons. Statistical analysis was performed with GraphPad Prism® 5.02 software. P values < 0.05 were considered significant.

RESULTS

Urodynamic evaluation of GK rats



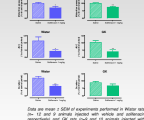
tative tracing of cystometrogram of conscious 18 weeks GK rat and age matched Wistar rat. Arrows represent micturition contractions. NVC: non-voiding contractions characteristic of detrusor overactivity.

Comparison of urodynamic parameters between Wistar and GK rats

	Wistar (n=21)	GK (n=20)
Micturition pressure (mmHg)	20.8 ± 0.8	23.7 ± 1.0 °
AUC micturition (mmHg x s)	114.6 ± 9.1	191.5 ± 13.4***
Duration micturition (s)	22.3 ± 0.9	27.4 ± 1.5 **
Threshold pressure (mmHg)	9.6 ± 0.6	8.8 ± 0.6
Basal pressure (mmHg)	5.9 ± 0.6	4.5 ± 0.4
Intercontraction interval (s)	443.7 ± 36.5	743.3 ± 58.2 ***
Bladder capacity (µI)	369.8 ± 30.5	619.4 ± 48.5 ***
Micturition volume (µI)	371.0 ± 44.1	636.9 ±65.1 **
Voiding efficiency (%)	97.6 ± 7.1	101.6±3.6
Amplitude of non-voiding contractions (mmHg)	3.1 ± 0.3	5.0 ± 0.4 ***
Frequency of non-voiding contractions (contractions per min)	0.3 ± 0.1	1.5 ± 0.2 ***
Volume threshold to elicit non-voiding contractions (%)	92.2 ± 1.6	65.2 ± 4.6 ***

◆The type 2 diabetes GK rat model displays severe diabetic bladder dysfunction characterized by bladder overactivity. They display increased micturition pressures, increased bla capacity and detrusor overactivity

characterizing

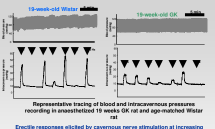


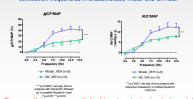
Data are mean ± SEM of experiments performed in Wistar rats
(n= 12 and 9 animals injected with vehicle and solifenacin
respectively) and GK rats (n=9 and 10 animals injected with
vehicle and solifenacin respectively).
'p-0.05." p=0.01 versus saling proup, Student's t-lest.

Solifenacin (1 mg/kg) inhibited the parameters characterizing the micturition

contraction in either GK or Wistar rats compared to saline injection without impacting voiding efficiency

Erectile function evaluation of GK rats





→ The erectile responses elicited by electrical stimulation of the cavernous nerve (6V, 1 ms for 45s) were considerably decreased in GK compared to Wistar rats at frequencies above 5 Hz

Effect of i.v sildenafil (0.3 mg/kg) or saline on erectile responses elicited by cavernous nerve stimulation at increasing stimulation frequencies in anaesthetized Wistar and GK rats





⇒Sildenafil (0.3 mg/kg) significantly increased the erectile response to ES CN in either Wistar or GK rats. The magnitude of this improvement was similar in both rats: at 15 Hz, AUC/MAP was increased by approximately 30% in both strains.

CONCLUSIONS

- The present study demonstrates that GK rats have many pathophysiological features in term of urological complications which are common to diabetic patients. They display diabetic bladder dysfunction characterized by detrusor overactivity, an increase in bladder capacity and micturition pressures. These rats also have erectile dysfunction associated.
- Furthermore, standard of care treatments for both disorders are effective in GK rats.
- Thus, GK rats represent a suitable and validated research model to better understand the pathophysiology of type 2 diabetes-associated bladder and erectile complications and to assess efficacy of new therapeutic agents targeting diabetic bladder and/or erectile dysfunctions.