BJPS

Brazilian Journal of Pharmaceutical Sciences vol. 49, n. 4, oct./dec., 2013

# Exendin-4 effects on islet volume and number in mouse pancreas

Layasadat Khorsandi\*, Fereshteh Nejad-Dehbashi

Cell and Molecular Research Center, Ahvaz Jundishapur University of Medical Sciences, Iran

The aim of this study was to evaluate Exendin-4 (EX-4) effects on islet volume and number in the mouse pancreas. Thirty-two healthy adult male NMRI mice were randomly divided into control and experimental groups. EX-4 was injected intraperitoneally (i. p.) at doses of 0.25 (E1 group), 0.5 (E2 group), and 1  $\mu$ g/kg (E3 group), twice a day for 7 consecutive days. One day after the final injection, the mice were sacrificed, and the pancreas from each animal dissected out, weighed, and fixed in 10% formalin for measurement of pancreas and islet volume, and determination of islet number by stereological assessments. There was a significant increase in the weight of pancreases in the E3 group. Islet and pancreas volumes in E1 and E2 groups were unchanged compared to the control group. The E3 group showed a significant increase in islet and pancreas volume (P < 0.05). There were no significant changes in the total number of islets in all three experimental groups. The results revealed that EX-4 increased pancreas and islet volume in non-diabetic mice. The increased total islet mass is probably caused by islet hypertrophy without the formation of additional islets.

**Uniterms:** Exendin-4/effects/experimental study. Pancreas/effects fo Exendin-4. Islet/hypertrophy. Stereology.

O objetivo deste estudo foi avaliar os efeitos do Exendin-4 (EX-4) sobre o volume e número de ilhotas no pâncreas. Trinta e dois camundongos NMRI machos saudáveis e adultos foram divididos ao acaso em grupos controle e grupos experimentais. EX-4 foi injetado intraperitonealmente (i. p.) nas doses de 0,25 (grupo E1), 0,5 (grupo E2) e 1 (grupo E3), duas vezes por dia durante 7 dias consecutivos. Um dia após a injeção final, os camundongos foram sacrificados e o pâncreas de cada animal foi dissecado, pesado e fixado em solução de formaldeído 10% para avaliação do volume do pâncreas e ilhotas e do número de ilhotas por métodos estereológicos. Observou-se aumento significativo no peso de pâncreas no grupo E3. O volume do pâncreas assim como das ilhotas não apresentou alterações nos grupos E1 e E2, quando comparados ao grupo controle No grupo E3 houve aumento significativo no volume do pâncreas e das ilhotas (P<0,05). Não se observaram alterações significativas no número de ilhotas nos três grupos experimentais. Os resultados revelaram que o EX-4 provoca aumento no volume total de ilhotas deve-se, provavelmente, a hipertrofia das ilhotas sem a formação de ilhotas adicionais.

Unitermos: Exendina-4/efeitos/estudo experimental. Pâncreas/efeito do Exendina-4. Ilhotas/hipertrofia. Estereologia.

# **INTRODUCTION**

Glucagon-like peptide-1 (GLP-1) is a peptide secreted from the gut in response to food. It acts directly on  $\beta$  cells, enhancing the effect of glucose in stimulating insulin secretion from these cells. When administered to diabetic mice, GLP-1 lowers blood glucose levels

and stimulates insulin secretion (Xu *et al.*, 1999). In addition, GLP-1 increases the  $\beta$ -cell mass by inducing the differentiation and neogenesis of ductal progenitor cells into islet endocrine cells (Hui *et al.*, 2001; Abraham *et al.*, 2002). In a previous *in vitro* study, was shown that GLP-1 is capable of enhancing fetal pig  $\beta$ -cell differentiation from progenitor epithelial cells as well as initiating their functional maturation in islet-like cell clusters (Hardikar *et al.*, 2002).

Exendin-4, a long-acting GLP-1 receptor (GLP-1R) agonist, binds to and activates the GLP-1R with the same

<sup>\*</sup>Correspondence: Layasadat Khorsandi. Cell and Molecular Research Center, Faculty of Medicine, Ahvaz Jundishapur University of Medical Sciences, P. O. Box: 61335, Ahvaz, Iran. E-mail: layasadat@yahoo.com

potency as GLP-1 (Xu *et al.*, 1999). Exendin-4 (EX-4) is resistant to the enzyme dipeptidyl peptidase 4 (DPT-IV), which is present in serum. GLP-1 is rapidly metabolized by DPP-IV (Kieffer *et al.*, 1995). It has been reported previously that EX-4 is capable of stimulating both the differentiation of  $\beta$  cells from ductal progenitor cells and proliferation of  $\beta$  cells when given to rats and human (Zhou *et al.*, 1999; Stoffers *et al.*, 2000; Kastin *et al.*, 2003).

Previous studies on EX-4 action were mostly performed in diabetic rodent models. However, some studies demonstrate that EX-4 has a beneficial effect in non-diabetic animals. It has been reported that EX-4 causes weight loss (De Fronzo *et al.*, 2005; Buse *et al.*, 2004; Kendall *et al.*, 2005). In the placebo-controlled component of the pivotal trials, which lasted 30 weeks, mean weight reduction ranged from between 1 and 3 kg compared with placebo. In open-label extensions, weight continued to decline over 2 years of treatment, by up to 5 kg from baseline (Blonde *et al.*, 2006; Buse *et al.*, 2007).

It is known that EX-4 can pass through the bloodbrain barrier (Kastin et al., 2003) and exert central effects, including promotion of neurotrophic or neuroprotective actions (Perry et al., 2000a; Perry et al., 2000b) and enhancement of cognitive functions (During et al., 2003). These findings suggest that GLP-1 receptor stimulation in the central nervous system plays a critical role in regulating neuronal plasticity and cell survival. Vella et al. (2003) reported that EX-4 and GLP-1 increase cortisol secretion in human subjects. However, neither of these alter insulin action in non-diabetic human subjects (Vella et al., 2003). Ranta et al. (2006) demonstrated that EX-4 protects against glucocorticoid-induced mouse beta-cells or INS-1 cell apoptosis. Chen et al. showed that EX-4 can inhibit rat cardiomyocyte apoptosis early after scald injury possibly by suppressing caspase-3 activity in the myocardium.

In spite of numerous experimental studies about EX-4 effects on various tissues in non-diabetic subjects, only one study has investigated its effects on the pancreas (Nachnani *et al.*, 2010). In the present study, the effect of EX-4 on islet volume and number in the mouse pancreas was investigated by using stereological procedures.

# MATERIAL AND METHODS

### Animals

In this study, 32 healthy adult male NMRI (Naval Medical Research Institute) mice (6–8 weeks old, 25-30 g) were used. The animals were obtained from Ahvaz

Jundishapur University of Medical Sciences, Experimental Research Center. This study was approved by the research ethics committee of Jundishapur University and carried out in an ethically proper way by following the guidelines provided. The animals were kept under standard laboratory conditions (12h-dark and 12 h-light cycle, relative humidity of  $50 \pm 5\%$  and  $22 \pm 3$  °C) for at least 1 week before the experiment and these conditions were maintained until the end of the experiment. Animal cages were kept clean, and commercial food (pellet) and water were provided *ad libitum*.

### **Experimental design**

The mice were randomly divided into control and experimental groups, all of which contained eight animals. EX-4 (Sigma) was dissolved in distilled water and injected intraperitoneally (i.p.) at doses of 0.25 (E1 group), 0.5 (E2 group), and 1  $\mu$ g/kg (E3 group), twice a day for 7 consecutive days. The dosage and duration of treatment with EX-4 was selected according to previous studies that demonstrated the beneficial effect of EX-4 on focal cerebral ischemia-induced infarction in rats (Briyal *et al.*, 2012). One day after the final injection, the mice were sacrificed by cervical dislocation, and the pancreas from each animal was dissected out, weighed, and fixed in 10% formalin.

### Stereological assessments

### Histology and sampling of sections

Each pancreas was embedded randomly in paraffin and sectioned exhaustively into  $5\mu$ m-thick sections. Figure 1 illustrates the sampling of sections. Three sections were collected onto each glass slide. With a random start point among the first 40 sections, every 40th section was sampled (the primary sections). In addition, two sections ahead of each primary section were sampled as the reference section. Because every section was 5 µm thick, it follows that there was 200 µm between the primary sections and 10 µm between a primary section and the corresponding reference section. All primary and reference sections were stained with hematoxylin and eosin (H & E).

### Microscopes and equipment

The sections were analyzed at 400× magnification by an MP3, Nr 3437 microscope (PZO, Poland) equipped with a projecting arm to project the image onto a monitor attached to the microscope. The applied probes used for the stereological examinations (point-counting grid or counting frame as described below) were superimposed



FIGURE 1- Sampling method of histological sections is shown.

onto the monitor so that the microscope projected the image onto the grid. We used two microscopes at the same time for counting the total number of islets in primary and reference sections.

#### Total volumes of islets and pancreas

Using step-lengths of 950 µm in the x-direction  $(\Delta x)$  and 750 µm in the y-direction  $(\Delta y)$ , all primary sections from each pancreas were systematically examined. A point-counting grid with 108 points, 1 of them encircled, was applied (Figure 2-A). Moving through all primary sections from the pancreata, the number of times 1 of the 108 points hit an islet was counted. An islet was defined as a cluster of cells with a minimum of three visible nuclei displaying the normal characteristics of islet endocrine cells (pale cytoplasm with approximately spherical nuclei). Simultaneously, the number of times the encircled point hit pancreatic tissue (exocrine pancreatic tissue, ducts, vessels, islets, etc.) was counted. The values for the total volume of pancreas and the islets of Langerhans were then calculated based on the Cavalieri principle (Gundersen et al., 1987; Bock et al., 2003).

1) V (pan) = 
$$a/p$$
 (pan) × N (p - p) × T ×  $\sum P$  (pan)  
= 0.1425 mm3 ×  $\sum P$  (pan)

where V (pan) is the total volume of pancreas, a/p(pan) is the area per point (in this case  $\Delta x \times \Delta y$  because only one point in the grid was used to count points that hit pancreas), N (p – p) is the number of sections between the primary sections (50 sections in this case), T is the section thickness (5 µm), and  $\sum P(pan)$  is the total number of points that hit pancreas.

2) V (isl) = 
$$a/p$$
 (isl) × N (p - p) × T ×  $\sum P$  (isl)  
= 1.319 × 10<sup>3</sup> mm<sup>3</sup> ×  $\sum P$  (isl)

where V (isl) is the total volume of islets, a/p (isl) is the area per point (in this case  $\Delta x \times \Delta y$ /99 because there were 99 points in the grid used to count points that hit islets), and  $\sum P$  (isl) is the total number of points that hit the islets.

Tissue shrinkage influences all stereologic size estimators whether distance, surface area, or volume. There is no exact unbiased way to obtain information about tissue deformation during tissue fixation and processing. The area of a piece of pancreas tissue before and after fixation/processing may be estimated, and the tissue shrinkage can be calculated as (Nyengaard, 1999):



**FIGURE 2** - Point counting grade (A) and unbiased counting frame (B).

#### Total number of islets

In another session, the sampling within the primary sections was performed, but an unbiased counting frame

(Figure 2-B) was attached to the monitor. The rules of the counting frame define objects completely outside the frame or objects that touch the exclusion lines (the full lines in Figure 2) as being outside the frame, whereas objects that are completely within the frame or touch only the inclusion lines (the dashed lines in Figure 2) are defined as being within the frame. We applied the dissector principle (Sterio, 1984) to count the islets. Whenever an islet profile was sampled by the counting frame, the corresponding position in the reference section was located with the other microscope, and it was then determined whether the islet was also visible in the reference section. An islet was counted if it appeared in the primary section but not in the reference section. Because the sampling of sections, as well as the within section sampling, were performed with known sampling fractions, the total number of islets can be calculated according to the fractionator principle (Bock et al., 2003; Gundersen et al., 1987) from:

$$N(isl) = \frac{N(p-p)}{N(p-r)} \times \frac{\Delta x \times \Delta y}{A(frame)} \times \Sigma Q^{-}(isl) = 34.53 \times \Sigma Q^{-}(isl)$$

where N (isl) is the total number of islets in the pancreas, N (p – p) is the number of sections between the primary sections, N (p – r) is the number of sections between a primary section and the corresponding reference section (two in this case),  $\Delta x$  and  $\Delta y$  are the step lengths, A(frame) is the area of the counting frame corrected for magnification (412.674 µm<sup>2</sup>), and  $\Sigma Q^{-}$  (isl) is the total number of islets counted in one pancreas (Bock *et al.*, 2003).

## RNA preparation and Reverse Transcription Polymerase Chain Reaction (RT-PCR)

Isolated pancreases were either used immediately or snap frozen in liquid nitrogen and stored at -80 °C until use. Using the RNeasy Mini kit (Qiagen), RNA was isolated from the tissues according to manufacturer's instructions. RT-PCR was performed using a One Step RT-PCR kit (Qiagen) which contains reverse transcriptase to synthesize cDNA from the RNA isolated and DNA polymerase for the PCR. RT-PCR conditions consisted of a 30 min step at 50 °C to allow the reverse transcriptase activity followed by 15 min at 95 °C to deactivate the reverse transcriptase and activate the Taq polymerase present in the enzyme mixture. The PCR process consisted of 6 s at 94 °C (denaturing step), 30 s at the annealing temperature (55 °C), and a 45 s step at 72 °C for extension, with all steps being repeated for 30 cycles. A final extension step lasted 10 min at 72 °C.

Primer sequences were as follows with the expected product length: Glut-2, sense 5' CA GCTGTCTCTGTGCTGCTGCTTGT3', antisense 5' GCCGTCATGCTCACATAACTCA3' (150 bp); Insulin, sense 5' TCTTCTACACACCCATGTCCC 3', antisense 5' GGTGCAGCACTGATCCAC 3', (149 bp); and GAPDH, sense 5' CTCTGGTGGACCTCATGGCCTAC 3', antisense 5' CAGCAACTGAGGGCCTCTCT 3' (105 bp) was used as the house keeping gene (Sun *et al.*, 2007).

### Statistical analysis

The data were analyzed using one-way ANOVA followed by the Post hoc LSD test and expressed as mean  $\pm$  SD. *P* < 0.05 was considered significant.

### RESULTS

As expected, mean body weight was equal in the four groups. Weight of pancreases in E1 and E2 groups were similar to the control group. There was a significant increase in the relative pancreas weight / body weight in the E3 group (Figure 3).



**FIGURE 3-** Relative pancreas weight / body weight in control and experimental groups. Values expressed as means  $\pm$  SD for 8 mice. \* P < 0.05 compared to control group.

The present study confirms a 30% tissue shrinkage in paraffin embedding. This shrinkage considered when the final results were reported.

In the E1 group, total islet volume and total pancreas volume were similar to the control group. Total islet numbers were also similar to the control group. Pancreas tissue showed normal architecture.

In the E2 group, total islet volume and total pancreas volume were slightly increased (P > 0.05). Total islet numbers were similar to the control group. No histopathological changes were observed in this group.

In the E3 group, total pancreas volume was significantly higher than the control group (P < 0.05). Total

islet volume was significantly increased compared to the control group (P < 0.05). Total islet numbers were similar to the control group. No histopathological changes were observed in this group. The results for total islet volume, total pancreas volume, and total islet numbers are depicted in Figure 4.



**FIGURE 4** - Total pancreas volume, total islet volume and total islet number of control and experimental groups. Values expressed as means  $\pm$  SD for 8 mice. \* P < 0.05 compared to control group.

To determine whether EX-4 affects  $\beta$ -cell islet function, the expression of Glut-2 and insulin genes were assessed using RT-PCR. As illustrated in Figure 5, high expression of Glut-2 and insulin was detected in EX-4treated mice. Expression of these genes was markedly higher in E2 and E3 groups compared to control and E1 groups.



**FIGURE 5** - The expression of insulin and Glut-2 genes in various groups by RT-PCR method is shown.

# DISCUSSION

Based on stereological methods (such as fractionator sampling and dissector counting), we found an increase in the total volume of islets in the experimental groups, whereas the total number of islets was equal in the four groups with a reasonably narrow confidence interval for the difference in means.

The importance of GLP-1 for stimulation of islet cell proliferation was originally demonstrated in lean 20-day-old normoglycemic mice (Edvell et al., 1999). Afterwards, several studies using in vivo models showed that GLP-1 can regulate islet growth mainly by controlling β-cell neogenesis (Xu et al., 1999; Stoffers et al., 2000; Perfetti et al., 2000; Tourrel et al., 2001; Tourrel et al., 2002). Park et al. showed that EX-4 and exercise promotes beta-cell function and mass in islets of diabetic rat. Xu et *al.* have also reported that EX-4 increases  $\beta$ -cell mass. Fan et al. (1999) reported that EX-4 improves blood glucose control in both young and aging normal non-diabetic mice. The authors showed that EX-4 treatment improved glycemic control in both 3-month and 20 to 22-month-old mice. In both groups of mice, the blood glucose lowering effect was independent of beta cell function as indicated by unchanged beta cell proliferation, insulin secretion or beta cell mass. However, high expression of Insulin 2 and Glut-2 genes in EX-4-treated cells was shown in the present study. In pancreatic  $\beta$ -cells, the glucose uptake is controlled by Glut-2, which is essential in the mechanism of glucose-induced insulin secretion (Olson et al., 1996). Glut-2 is the glucose sensor of  $\beta$  cells that leads to the production of insulin (Lopes Da Costa et al., 2004).

GLP-1 increases insulin secretion and the biosynthesis of important  $\beta$ -cell products besides insulin: glucokinase and Glut-2 glucose transporters (Verspohl *et al.*, 2009). The increase in expression of these genes probably induces abnormally elevated secretion of insulin and causes hypoglycemia in non-diabetic animals.

As mentioned above, in this study the volume of islets was higher in EX-4-treated mice. One mechanism responsible for the expansion of islet mass is inhibition of apoptosis (Chen *et al.*, 2011; Farilla *et al.*, 2003; Kwon *et al.*, 2009). It has also been shown that human islets treated with GLP-1 have a down-regulation of caspase-3 at the levels of mRNA of the active protein and up-regulation of the anti-apoptotic protein Bcl-2 (Farilla *et al.*, 2003). A second mechanism responsible for the expansion of  $\beta$ -cell mass is enhanced cell proliferation or neogenesis.

Tourrel *et al.*, by using a recognized model of  $\beta$ -cells regeneration (neonatalWistar rats injected with streptozotocin, so-called n0-STZ), showed that GLP-1 and Exendin-4, applied during the neonatal period, strongly stimulated  $\beta$ -cell regeneration mainly by  $\beta$ -cell neogenesis (Tourrel *et al.*, 2001). Furthermore, treatment of diabetic Goto-Kakizaki (GK) rats with GLP-1 or Exendin-4 from day 2 to day 6 after birth resulted in stimulation of  $\beta$ -cell neogenesis and proliferation with persistent expansion of  $\beta$ -cell mass detected at adult age (Tourrel *et al.*, 2002). However, the present study revealed that EX-4 caused no change in the number of islets. This indicates that EX-4 has no neogenesis effects on islet cells in non-diabetic adult animals.

It has been stated in a literature review that new islets do develop under certain experimental conditions, such as after partial pancreatectomy, where the formation of new islets has been clearly demonstrated (Nachnani *et al.*, 2010). Other anatomical structures, such as kidney glomeruli, also lack the ability of hyperplasia and instead become hypertrophic with an increased demand, probably because of the highly specific structure of the neurovascular and tubular systems necessary for appropriate function. Possibly, the architecture (i.e., the intra-islet vascular structure) of the islets is complex to such a degree that it only allows new islets to be formed during the formation, growth, or regeneration of the pancreas during fetal life or after partial pancreatectomy (Bock *et al.*, 2003).

Nachnani *et al.* evaluated the histological and biochemical effects of EX-4 on the pancreas in rats. They showed that animals treated with Exendin-4 had pancreatic acinar inflammation, pyknotic nuclei and weighed significantly less than control rats. However, in the present study no evidence of pancreatic acinar inflammation or histopathological changes were observed.

# CONCLUSION

In this study, we demonstrated that EX-4 increased pancreas and islet volume in non-diabetic mice. The increased total islet mass is probably caused by islet hypertrophia without the formation of additional islets. This study also revealed that EX-4 can enhance the expression of insulin and Glut-2 genes, where this may induce hypoglycemia in non-diabetic mice. Further experiments are needed to clarify the exact mechanism of islet hypertrophy induced by EX-4 and other GLP-1 agonists.

## ACKNOWLEDGEMENT

This research was supported by a Grant (CM-004) from the research council of the Ahvaz Jundishapur University of Medical Sciences in 2011.

# REFERENCES

- ABRAHAM, E.J.; LEECH, C.A.; LIN, J.C.; ZULEWSKI, H.; HABNER, J.F. Insulinotropic hormone glucagon-like peptide-1 differentiation of human pancreatic islet-derived progenitor cells into insulin producing cells. *Endocrinology*, v.143, p.3152-3161, 2002.
- BLONDE, L.; KLEIN, E.J.; HAN, J.; ZHANG, B.; MAC, S.M.; POON, T.H.; TAYLOR, K.L.; TRAUTMANN, M.E.; KIM, D.D.; KENDALL, D.M. Interim analysis of the effects of exenatide treatment on A1C, weight and cardiovascular risk factors over 82 weeks in 314 overweight patients with type 2 diabetes. *Diabetes Obes. Metab.*, v.8, p.436-447, 2006.
- BOCK, T.; PAKKENBERG, B.; BUSCHARD, B.; BUSCHARD, K. Inctrased islet volume but unchanged islet number in *ob/ob* mice. *Diabetes*, v.52, p.1716-1722, 2003.
- BRIYAL, S.; GULATI, K.; GULATI, A. Repeated administration of exendin-4 reduces focal cerebral ischemia-induced infarction in rats. *Brain Res.*, v.1427, p.23-34, 2012.
- BUSE, J.B.; HENRY, R.R.; HAN, J.; KIM, D.D.; FINEMAN, M.S.; BARON, A.D. Effects of exenatide (exendin-4) on glycemic control over 30 weeks in sulfonylureatreated patients with type 2 diabetes. *Diabetes Care*, v.27, p.2628-2635, 2004.

- BUSE, J.B.; KLONOFF, D.C.; NIELSEN, L.L.; GUAN, X.; BOWLUS, C.L.; HOLCOMBE, J.H.; MAGGS, D.G.; WINTLE, M.E. Metabolic effects of two years of exenatide treatment on diabetes, obesity, and hepatic biomarkers in patients with type 2 diabetes. *Clin Ther.*, v.29, p.139-153, 2007.
- CHEN, Y.H.; WANG, J.H.; LI, Z.Q.; YI, Z.H. Effects of exendin-4 on rat cardiomyocyte apoptosis early after severe scald injury. *Nan. Fang. Yi. Ke. Da Xue. Xue. Bao.*, v.31, p.1101-1104, 2011.
- DE FRONZO, R.A.; RANTER, R.E.; HAN, J.; KIM, D.D.; FINEMAN, M.S.; BARON, A.D. Effects of exenatide (exendin-4) on glycemic control and weight over 30 weeks in metformin-treated patients with type 2 diabetes. *Diabetes Care*, v. 28, p.1092-1100, 2005.
- DURING, M.J.; CAO, L.; ZUZGA, D.S.; FRANCIS, J.S.; FITZIMONS, H.L.; JIAO, X.; BLAND, R.J.; KLUGMANN, M.; BLANKS, W.A.; DRUCKER, D.J.; HAILE, C.N. Glucagon-like peptide-1 receptor is involved in learning and neuroprotection. *Nat. Med.*, v.9, p.1173-1179, 2003.
- EDWEL, A.; LINDSTROM, P.; EDVELL, A.; LINDSTROM, P. Initiation of increased pancreatic islet growth in young normoglycemic mice (Umea +/?). *Endocrinology*, v.140, p.778-783, 1999.
- FAN, R.; KANG, Z.; HE, L.; CHAN, J.; XU, G. Exendin-4 improves blood glucose control in both young and aging normal non-diabetic mice, possible contribution of beta cell independent effects. *PloS One*, v.6, p.e20443, 2011.
- FARILLA, L.; BULOTTA, A.; HIRSHBERG, B.; LI-CALAZI, S.; KHOURY, N.; NOUSHMEHR, H.; BETOLOTTO, C.; DI MARIO, U.; HARLAN, D.M.; PERFETTI, R. Glucagon-like peptide1 inhibits cell apoptosis and improves glucose responsiveness of freshly isolated human islets. *Endocrinology*, v.144, p.5149-5158, 2003.
- GUNDERSEN, H.J. Stereology of arbitrary particles: are view of unbiased number and size estimators and the presentation of some new ones, in memory of WilliamR. Thompson. J. *Microsc.*, v.143, n.1, p.3-45, 1986.
- GUNDERSEN, H.J.; JENSEN, E.B. The efficiency of systematic sampling in stereology and its prediction. J. *Microsc.*, v.14, p.229-263, 1987.

- HARDIKAR, A.A.; WANG, X.Y.; WILLIAMS, L.; KWOK, J.; WONG, R.; YAO, M.; TUCH, B.E. Functional maturation of fetal porcine beta cells by glucagon-like peptide 1 and cholecystokinin. *Endocrinology*, v.143, p.3505-3514, 2002.
- HUI, R.; WRIGHT, C.; PERFETTI, R. Glucagon-like peptide 1 induces differentiation of islet duodenal homeobox-1 positive pancreatic ductal cells into insulin-secretion cells. *Diabetes*, v.50, p.785-796, 2001.
- KASTIN, A.J.; AKERSTROM, V. Entry of exendin-4 into brain is rapid but may be limited at high doses. *Int. J. Obes. Relat. Metab. Disord.*, v.27, p.313-318, 2003.
- KENDALL, D.M.; RIDDLE, M.C.; ROSENSTOCK, J.; ZHUANG, D.; KIM, D.D.; FINEMAN, M.S.; BARON, A.D. Effects of exenatide (exendin-4) on glycemic control over 30 weeks in patients with type 2 diabetes treated with metformin and a sulfonylurea. *Diabetes Care*, v.28, p.1083-1091, 2005.
- KIEFFER, T.J.; MCINTOSH, C.H.; PEDERSON, R.A. Degradation of glucose-dependent insulinotropic polypeptide and truncated glucagon-like peptide-1 *in vitro* and *invivo* by dipeptidyl peptidase IV. *Endocrinology*, v.136, p.3585-3596, 1995.
- LOPES DA COSTA, C.; SAMPAIO DE FREITAS, M.; SANCHES MOURA, A. Insulin secretion and GLUT-2 expression in undernourished neonate rats. *J. Nutr. Biochem.*, v.15, p.236-241, 2004.
- NACHNANI, J.S.; BULCHANDANI, D.G.; NOOKALA, A.; HERNDON, B.; MOLTENI, A.; PANDYA, P.; TAYLOR, R.; QUINN, T.; WEIDE, L.; ALBA, L.M. Biochemical and histological effects of exendin-4 (exenatide) on the rat pancreas. *Diabetologia*, v.53, p.153-159, 2010.
- NYENGAARD, J.R. Stereologic methods and their application in kidney research. *J. Am. Soc. Nephrol.*, v.10, p.1100-1123, 1999.
- OLSON, A.L.; PESSIN, J.E. Structure function, and regulation of the mammalian facilitative glucose transporter gene family. *Annu. Rev. Nutr.*, v.16, p.235-256, 1996.
- PERFETTI, R.; ZHOU, J.; DOYLE, M.E.; EGAN, J. M. Glucagon-like peptide-1 induces cell proliferation and pancreatic-duodenum homeobox-1 expression and increases endocrine cell mass in the pancreas of old, glucose-intolerant rats. *Endocrinology*, v.141, p.4600-4605, 2000.

- PERRY, T.; HAUGHEY, N.J.; MATTSON, M.P.; EGAN, J.M.; GRIEG, N.H. Protection and reversal of excitotoxic neuronal damage by glucagon-like peptide-1 and exendin-4. *J. Pharmacol. Exp. Ther.*, v.302, p.881-888, 2002a.
- PERRY, T.; LAHIRI, D.K.; CHEN, D.; ZHOU, J.; SHAW, K.T.; EGAN, J.M.; GRIEG, N.H. A novel neurotrophic property of glucagon-like peptide 1: a promoter of nerve growth factor-mediated differentiation in PC12 cells. *J. Pharmacol. Exp. Ther.*, v.300, p. 958-966, 2002b.
- PARK, S.; HONG, S.M.; SUNG, S.R. Exendin-4 and exercise promotes beta-cell function and mass through IRS2 induction in islets of diabetic rats. *Life Sci.*, v.82, p.9-10, p.503-511, 2008.
- RANTA, F.; AYRAM, D.; BERCHTOLD, S.; ADKINS, A.S.; BASU, R.; RIZZA, R.A. Dexamethasone induces cell death in insulin-secreting cells, an effect reversed by exendin-4. *Diabetes*, v.55, p.1380-1390, 2006.
- STOFFERS, D.A.; KIEFFER, T.J.; HUSSAIN, M.A.; DRUCKER, D.J.; BONNER-WEIR, S.; HABENER, J.F.; EGAN, J.M. Insulinotropic glucagon-like peptide 1 agonists stimulate expression of homeodomain protein IDX-1 and increase islet size in mouse pancreas. *Diabetes*, v.49, p.741-748, 2004.
- SUN, Y.; ZHANG, L.; GU, H.F.; HAN, W.; REN, M.; WANG, F.; GONG, B.; WANG, L.; GUO, H.; XIN, W.; ZHAO, J.; GAO, L. Peroxisome proliferator-activated receptor-1 regulates the expression of pancreatic/duodenal homeobox-1 in rat Insulinoma (INS-1) cells and ameliorates glucose-induced insulin secretion impaired by palmitate. *Endocrinology*, v.149, p.662-671, 2007.
- TOURREL, C.; BAILBE, D.; LACOME, M.; MEILE, M.J.; KERGOAT, M.; PORTHA, B. Persistent improvement of type 2 diabetes in the Goto-Kakizaki rat model by expansion of the beta-cell mass during the prediabetic period with glucagon-like peptide-1 or exendin-4. *Diabetes*, v.51, p.1443-1452, 2002.

- TOURREL, C.; BAILBE, D.; MEILE, M.J.; KERGOAT, M.; PORTHA, B. Glucagon-like peptide-1 and exendin-4 stimulate beta cell neogenesis in streptozotocin-treated newborn rats resulting in persistently improved glucose homeostasis at adult age. *Diabetes*, v.50, p.1562-1570, 2001.
- VELLA, A.; SHAH, P.; REED, A.S.; ADKINS, A.S.; BASU, R.; RIZZA, R.A. Lack of effect of exendin-4 and glucagon-like peptide-1-(7,36)-amide on insulin action in non-diabetic humans. *Diabetologia*, v.46, p.1589, 2003.
- VERSPOHL, E.J. Novel therapeutics for type 2 diabetes: Incretin hormone mimetics (glucagon-like peptide-1 receptor agonists) and dipeptidyl peptidase-4 inhibitors. *Pharmacol. Ther.*, v.124, p.113-138, 2009.
- XU, G.; STOFFERS, D.A.; HABENER, J.F.; BONNER-WEIR, S. Exendin-4 stimulates both b-Cell replication and neogenesis, resulting in increased b-Cell mass and improved glucose tolerance in diabetic rats. *Diabetologia*, v.48, p.2270-2276, 1999.
- ZHOU, J.; WANG, X.; PINEYRO, M.A.; EGAN, J.M.; ZHOU, J.; WANG, Y.; PINEYRO, M.A.; EGAN, J.M. Glucagonlike peptide 1 and exendin-4 convert pancreatic AR42J cells into glucagon- and insulin-producing cells. *Diabetes*, v.48, p.2358-2366, 1999.

Received for publication on 05<sup>th</sup> August 2012 Accepted for publication on 29<sup>th</sup> November 2012