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Amylin induces hypoglycemia in mice

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ABSTRACT

Amylin is a 37-aminoacid pancreatic protein that exerts control over several metabolic events such as glycemia and lacticemia. Amylin has long been shown to induce increases in arterial plasma glucose. We decided to investigate whether amylin plays additional roles in the glucose metabolism. We evaluated glucose homeostasis using whole blood from the tail tip of fasting, conscious, unrestrained normal and streptozotocyn-induced diabetic mice following subcutaneous administration of mouse amylin. Subcutaneous injection of 1 µg mouse amylin caused a transient decrease in whole blood glucose in both normal and diabetic mice in the absence of insulin. The blood glucose levels were lowest approximately 2 hours after amylin administration, after that they gradually recovered to the levels of the control group. The hypoglycemic effect followed a dose-dependent response ranging from 0.1 to 50 µg / mouse. These results reveal the ability for amylin in the direct control of glycemia at low doses in the absence of insulin.

Key words: amylin, islet associated polypeptide, glycemia, diabetes, mice, hypoglycemia.

INTRODUCTION

Amylin, also known as islet associated / amyloid polypeptide (IAPP), is a 37-residue protein which is carboxy-amidated and contains a disulfide bridge between cysteines Cys2 and Cys7. Amylin is stored and co-secreted with insulin (Hartter et al. 1991) in response to various metabolic conditions. Plasma amylin concentrations depends on the body’s metabolic conditions and are typically in the range of 2 to 20 pM in fasting and fed conditions, respectively (Butler et al. 1990), and at reduced levels or entirely absent in diabetic individuals (Hartter et al. 1990).

Soon after the discovery of amylin as a pancreatic deposit (Cooper et al. 1987, Westermark et al. 1987), the initial attempts to elucidate its effects on carbohydrate metabolism failed. The limited solubility of human amylin in aqueous solution results in the lack of biological activity, whereas water-soluble mouse/rat (same sequence) amylin does exhibit bioactivity (Bretherton-Watt et al. 1990, Balasubramaniam et al. 1991, Young et al. 1991).

Subsequent studies using high amylin doses and anesthetized animal models allowed the collection of relatively large arterial blood volumes and
enabled a greater variety of biochemical analyses. Since then, amylin has been found to be involved in the regulation of glycemia, lacticemia, food intake, gastric emptying and the secretion of insulin and glucagon (Young 2005). The hyperglycemia induced by subcutaneous or intravenous administration of amylin is preceded by an increase in plasma lactate and has been observed in the arterial blood of both fasting and fed anesthetized rats and mice (Wang et al. 1991, 1992, Young et al. 1991, 1996). This hyperglycemia is attributed to the combined, dose-dependent effects of amylin’s inhibition of insulin and glucagon secretion and its stimulation of gluconeogenesis (Young 2005). One study also found a greater effect of amylin on the liver compared with the peripheral tissues (Koopmans et al. 1991).

These effects of amylin on arterial blood glucose and other metabolites have been reported in the literature for over two decades. However, most data have resulted from measurements of the arterial blood of anesthetized animals. Halothane, isoflurane and other common anesthetic compounds have systemic metabolic effects including decreasing insulin secretion and causing hyperglycemia, hypocalcemia and hyperlactemia (Bito and Eakins 1969, Young et al. 1991, Wang et al. 1991, Farrokhnia et al. 2009, Zuurbier et al. 2008). Therefore, they might interfere in the measurement of discrete metabolic and physiological events.

In this context, we decided to investigate whether amylin exerts any additional roles in glucose metabolism. To address this issue, we evaluated the effects of mouse amylin on glycemia in fasting, conscious unrestrained mice by monitoring the whole blood glycemia in mice tail tips (Ayala et al. 2010). We report here a dose-dependent, transient decrease in blood glucose induced by subcutaneous mouse amylin. These data introduce a new physiological role for amylin, which in conjunction with the well-known hyperglycemic effects of amylin, highlights the importance of its participation in the tight hormonal regulation of homeostasis.

**MATERIALS AND METHODS**

**REAGENTS**

Mouse amylin (CAS 122384-88-7) was obtained from GenScript (cat. # RP11280, lot #55613-1 and #128567001040611LQ; mouse amylin is identical in sequence to rat amylin). Distilled water was deionized to less than 1.0 μS and filtered through a 0.22μm-pore membrane in a water purification system prior to use. All other reagents were of analytical grade. All buffers and solutions were prepared immediately prior use.

**PHARMACOLOGICAL EVALUATION OF AMYLIN**

Eight-weeks-old Swiss male mice (27 g ± 1 g) were divided into four major groups: normal (both control and amylin) and diabetic (both control and amylin). The animals were housed in a temperature-controlled room with a 12:12 hours light-dark cycle. Type 1 diabetes was induced by one intraperitoneal (ip.) injection of STZ (200 mg/kg) dissolved in fresh citrate buffer (100 mM, pH 4.5) (Da et al. 2010, Wu and Huan 2008, Xu et al. 2006, Ogawa et al. 1990). Normal (non-diabetic) group received just the vehicle (0.1mL). After STZ administration (5 days), blood was drawn from mice by tail snip and glucose level was evaluated as described below. Diabetes was defined as blood glucose concentration higher than 300 mg/dL. Water and food were available ad libitum, and food was suspended 4 h before the experiments. All mice were fasted throughout the experiments and kept at constant temperature. The first group received either 100 μL of saline (0.9 % NaCl) or phosphate buffered saline (8.1 mM Na2HPO4, 1.8 mM KH2PO4, 2.7 mM KCl, 137 mM NaCl, pH 7.4) as described in the figure legends. The second group received 100 μL of the test solutions at varying doses. Solutions were administrated by subcutaneous injection using a standard 29-gauge needle. Glucose concentrations were monitored by using whole blood from the tail tip of the conscious, unrestrained mice using pre-calibrated glucometers.
HYPOGLYCEMIC ACTIVITY OF AMYLIN

(Accu-Chek® Active, Roche Diagnostics, Germany; Serial Nos GC02476995 and GN08146937) before injection and at repeated intervals after administration for up to 20 hours. This protocol is in accordance with the recommendations of the Mouse Metabolic Phenotyping Center Consortium (MMPC) from the National Institutes of Health (NIH) for animal experimentation and for evaluating the effects of metabolic compounds on glucose homeostasis (Ayala et al. 2010), and followed the ‘Principles of laboratory animal care’ (NIH publication no. 85–23, revised 1985). A control performance test was conducted by evaluating of the insulin-induced hypoglycemia and glucose-induced hyperglycemia using the same experimental protocol (not shown). Amylin activity was confirmed by evaluating the hyperglycemic effect of 100 μg mouse (=rat) amylin (subcutaneous) on rats (approx. weight 370 g) fasted for 5 hours and anesthetized with halothane using arterial blood taken from the carotid artery (Wang et al. 1991, Young et al. 1991). This study was approved by the Institutional Bioethics Committee on Animal Care and Experimentation (IBCACE) at UFRJ.

STATISTICAL ANALYSIS

A paired t-test was used to compare the differences in pharmacological response. Values of *p*<0.05 were considered significant. The analyses were performed using SigmaStat within SigmaPlot 11 (Systat Software Inc).

RESULTS

To understand the effects of amylin on glucose metabolism, we have monitored the glycemic effects of murine amylin in mice. Swiss male mice were fasted for 4 hours before subcutaneous administration of 100 μL of either saline or 1 μg mouse amylin in saline, and changes in glycemia were monitored at given intervals with blood taken from the tail tip of conscious, unrestrained mice (Fig. 1). Amylin resulted in the depression of the whole blood glucose levels in normal (i.e., non-diabetic) mice for the test group compared to the control group receiving saline only within the first 2 hours after administration (Fig. 1A). Approximately 6 hours after treatment the glycemia of both groups had converged to similar levels (Fig. 1A).

We have further evaluated the effect of amylin over the glycemia of streptozotocin-induced diabetic mice (Da et al. 2010, Wu and Huan 2008, Xu et al. 2006, Ogawa et al. 1990). Amylin was administrated through subcutaneous route and glycemia was monitored from whole blood. Amylin administration resulted in a rapid reduction of blood glucose within a few hours followed by a progressive restoration of glycemia to similar level of the control group, i.e., streptozotocin-induced diabetic mice receiving saline (Fig. 1B).

Figure 1 - Amylin induces hypoglycemia in fasting mice. Either a control solution (■, 0.9 % saline) or mouse amylin (□, 1 μg in 0.9 % saline) was administered via subcutaneous injection in 4 h-fasted mice (n = 4 per group), both A) normal (non-diabetic) and B) streptozotocin-induced type-1 diabetic mice. Glycemia was monitored using blood collected from the tail tip while the mice were fasting, conscious and unrestrained. Hypoglycemia was observed in the amylin-treated group (*p*=0.008; 2h, *p*=0.05). Symbols represent mean ± s.e.m. Further details can be found in the Methods section.

We further analyzed the relative changes in glycemic values induced by amylin. The normalized glycemic values showed a transient behavior, whereby the maximum decline in blood glucose occurred approximately 1 to 2 hours after amylin administration, followed by a progressive recovery of glycemic values to normal levels (Fig. 2). These results were reproducible regardless of
room temperature (19 or 24 °C), amylin solution composition (0.9 % NaCl saline or PBS buffer), analyst, glucometer equipment, amylin batch or the origin (vivarium) of the mice (Fig. 2).

**Figure 2 - Time-dependence on the hypoglycemia response to subcutaneous mouse amylin.** Mice were treated with either amylin (1 μg in either saline or PBS solution) or a control solution of either saline or PBS administered via subcutaneous injection in 4 h-fasted mice (n≥3 per group). Glycemia was monitored using blood collected from the tail tip while the mice were fasting, conscious and unrestrained. The results from the treatment group were subtracted from the control group, and the results are expressed here as percentage change. Experiments were performed by different analysts at 25 °C (●) or 19 °C (■), by using either 0.9 % saline (■) or PBS (●) as the amylin diluting solution, using different glucometers, amylin batch and animals from distinct vivarium. Further details can be found in the Methods section.

We further evaluated the effects of varying the amylin dose on glycemia (Fig. 3). At very low doses (0.01 μg/mouse) there was no observable effect compared to the control. Higher doses led to a progressive decrease in glycemic values, achieving a level approximately 80% of that of the control mice. In summary, a small but sensitive ability to decrease glycemic values was observed with doses from 0.1 to approximately 50 μg amylin / mouse.

**DISCUSSION**

Amylin physiology and pharmacology has been studied intensely for two decades by independent laboratories. Amylin induces hyperglycemia in vivo regardless of the model studied, rat, mouse or human, in both normal and diabetic individuals. To date, hypoglycemic effects have not been reported, even at supraphysiological doses of amylin (Nyholm et al. 1996).

In the present study, we evaluated the hypoglycemic effects of subcutaneous mouse amylin using whole blood collected from the cut tail tip of mice. The observed hypoglycemic effects occurred over a small concentration range, with the maximum effect at 1.0 μg/mouse, decreasing at higher doses. This effect was not restricted to murine amylin, since a hypoglycemic effect in mice was also observed using human amylin released from polymeric nanoparticles (Guerreiro et al. 2012) and from phosphatidylcholine liposomes using the same experimental pharmacology protocol reported here. These results provide further evidences that the hypoglycemic effects are not isolated attributes of mouse amylin or a device or formulation concern, but are most likely a general, physiological response to circulating amylin. We have also shown here that type
1-diabetic mice were fully responsive to amylin in the absence of insulin, suggesting a direct role of amylin in controlling glycemia.

Our present findings do not contradict the hyperglycemic effects reported previously. Instead, we consider that the hypoglycemia reported here is peripheral and occurs to a small but significant extent in a concentration-dependent manner. This concentration dependence occurs over a different range from that observed for hyperglycemia in arterial plasma; the latter occurs over a large extent, making it difficult to observe a discreet, peripheral hypoglycemic effect.

Collectively, these data suggest that amylin-induced hyperglycemia and hypoglycemia events are both reliable, dose-dependent effects, exerting their effect at both physiologic (low concentration) and pharmacologic (high concentration) conditions. Furthermore, they also suggest the existence of a relationship between local trends in glucose homeostasis and the different activities of amylin at varying sites, such as specific organs, tissues, interaction and signaling mechanism, at distinct concentration ranges. The hypoglycemic effect reported here introduces a new physiological and pharmacological action of amylin in living individuals. Further studies are needed to advance the understanding on how amylin affects biological systems.

CONCLUSIONS

Amylin is a pancreatic hormone known for acting over a large series of physiologic phenomena, including the regulation of glucose homeostasis, and is absent or present in reduced level in diabetes. In the present study we have shown that amylin can directly induces hypoglycemia in fasting, conscious unrestrained mice, in a dose-dependent hypoglycemic effect of amylin. Data support a new physiologic activity for amylin. We do believe that both the hypoglycemic and hyperglycemic effects of amylin are of equal importance, thought clear separated events regarding tissue specificity, physiopathologic conditions, and depend on the concentration range for the hormone at the site of action. Our data provides a landmark view on new physiologic functions for such an important hormone, and may open new perspectives in the study of system biology involving amylin.

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RESUMO

Amilina é uma proteína de 37 aminoácidos que exerce controle sobre diversos eventos metabólicos, tais como glicemia e lacticemia. Tem sido mostrado que amilina induz aumento da glicemia plasmática arterial. Nós decidimos investigar se amilina exerce função adicional na regulação do metabolismo de glicose. Nós avaliámos a homeostase de glicose empregando sangue total da ponta da cauda de camundongos normais e diabéticos em condição livre de anestésico e não-confinados. Injeção subcutânea de 1 μg por camundongo causou o decréscimo transiente na glicemia tanto em camundongos normais quanto em diabéticos na ausência de administração de insulina. Os níveis de glicose atingiram o mínimo em aproximadamente 2 horas após administração de
amilóide, glicemia, diabetes, camundongos, hipoglicemia.

Palavras-chave: amilina, polipeptídeo pancreático amilóide, glicemia, diabetes, camundongos, hipoglicemia.

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