Characterization of the *Bombyx mori* homolog of islet homeostasis protein (IHoP) cDNA: *BmIHoP* controls insulin secretion from the β-cells.

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**Abstract**

Islet homeostasis protein (IHoP) was firstly characterized as a novel protein that regulates glucagon synthesis in alpha-cells of pancreatic islet. Here, we report a cDNA designed *BmIHoP* encoding an IHop homolog cloned from the fat body of the silkworm, *Bombyx mori*, using conserved domain homology search method. The resultant *BmIHoP* translated into a protein that encodes 693 amino acids, with a theoretical isoelectric point of 5.15 and a molecular weight of 77.5 kDa. Homology analysis revealed that *BmIHoP* has higher similarity at the amino acid level with those of other species, 52% identity with *H. sapiens*, 52% with *M. musculus*, 51% with *D. rerio*, and 73% with *D. melanogaster*. The expression of *BmIHoP* was detected in all tissues tested with 4 fold higher expression in the midgut, Malpighian, ovary and testis, and stronger expression in the larval fat body at 5th-3rd instar. The overexpression of *BmIHoP* in the β-cells reduced insulin secretion, which is not associated with SOX17 regulation. To the best of our knowledge, this is the first report of *B. mori* IHoP cDNA, *BmIHoP*, and its associated role in controlling insulin secretion from the β-cells.

**Keywords:** Silkworm, *Bombyx mori*, Islet homeostasis protein, cDNA, Insulin.

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**Introduction**

The pancreatic islets are small islands composed of 5 different types of cells that produce peptide hormones including glucagon by alpha cells (α-cells), insulin by beta cells (β-cells), somatostatin by delta cells (δ-cells), pancreatic polypeptide by gamma cells (PP-cells), and ghrelin also known as lenomorelin by epsilon cells (ε-cells) [1-3]. The synthesized hormones are secreted directly into the bloodstream for the homeostasis of blood glucose, fatty acids, triglycerides, and amino acids [4]. Especially, maintaining homeostasis of blood glucose level is regulated by two kinds of hormones, insulin, and glucagon. It is important to maintain the balance between insulin and glucagon in blood, and the involved mechanism is described in brief as follows: when blood glucose level is high, stimulation of pancreas takes place with the subsequent release of insulin from β-cells. In the case of glucagon, the hormone is released from α-cells of the pancreas, when blood glucose levels are low [5,6].

The liver is the main target organ of glucagon, which controls glucose secretion from the stored liver glucose. Glucagon is biosynthesized from the cleavage of proglucagon by proprotein convertase 2 in pancreatic islet α-cells [7,8]. While secretion of glucagon is stimulated by hypoglycemia, epinephrine, arginine, alanine, alanine transaminase, acetylcholine, and cholecystokinin, glucagon secretion is inhibited by somatostatin, insulin, PPARγ/retinoid X receptor heterodimer, increased free fatty acids, and urea production [9-11].

Although extensive research has been carried out with emphasis on type-1 diabetes, the underlying acute mechanisms and pathogenesis are not completely understood. Further research is required to gain deeper insight into the diagnosis and prevention of type-1 diabetes. Recently, a study reported that a gene co-localized with α-cells and encoding islet homeostasis protein (IHoP) produces glucagon from the pancreatic Langerhans islets. The authors demonstrated that IHoP strongly associates with glucagon synthesis, which exhibits its effects on insulin regulation. Inhibition of IHoP expression resulted in the loss of glucagon expression, which induced low blood glucose levels and cellular apoptosis [12]. The data suggested that IHoP might be actively involved in maintaining pancreas homeostasis through the regulation of α-cells and β-cells.
already reported homologous genes. Also, we demonstrate that sequence analysis, expression patterns of tissues and BmIHoP
supplements or ingredients may play few important roles in practically used, an oral toluene method that avoids digestion developmental stages; comparative evolutionary analyses of silkworm in the treatment of diabetes at the experimental level [14-16]. Henceforth, we can consider that BmIHoP as a supplements or ingredients may play few important roles in maintaining pancreas homeostasis in the mammalian. To be practically used, an oral toluene method that avoids digestion in the stomach and gut into peptides should be considered [17]. In the present work, we have isolated a cDNA encoding BmIHoP from silkworm fat body and characterized by sequence analysis, expression patterns of tissues and developmental stages; comparative evolutionary analyses already reported homologous genes. Also, we demonstrate that BmIHoP expression in the β-cells reduced insulin secretion in which SOX17 expression was not related and induced apoptosis [18].

Materials and Methods

Cloning of BmIHoP cDNA

Total RNA was extracted from the fat body of B. mori (Jam123 × Jam124) using Trizol reagent (Invitrogen, Carlsbad, CA, USA) and was further treated with RNase-free DNase-I to remove any potential genomic DNA contamination. Candidate DNA fragments were amplified and subsequently confirmed via direct DNA sequencing. The BLAST search was used for the gene identification. PCR was performed using the primers designed with Primer3 (http://simgene.com/Primer3) based on ‘Conserved Domain Databases’ (NCBI, National Center for Biotechnology Information) and ‘Motif Databases’ (GenomeNet, Institute for Chemical Research, Kyoto University, Japan), and the number of PCR cycles was optimized in order to stay within the linear range of amplification. The primers used were as follows: BmIHoP-FOR (5′-CTG TCC ATG GCC AAC ATT TCA-3′) and BmIHoP-REV (5′-CCA TCT TAA CTT CCA AAT CGA TTC-3′). The resulting PCR product was sub-cloned in the TOPO TA Cloning® plasmid (Invitrogen) and sequenced to confirm specificity. Several clones were finally sequenced to confirm of some mutations may occur by the DNA polymerase in the cloned cDNAs. All other chemicals and drugs were purchased from Sigma Chemical (St. Louis, MO).

Characterization of BmIHoP

An open reading frame (ORF), molecular weight, and theoretical isoelectric point were estimated by ExPASy (http://www.expasy.org). The multiple protein sequence alignments and phylogenetic evolutionary analyses were generated using NCBI (http://www.ncbi.nlm.nih.gov/).

Gene expression analysis by RT-PCR

Silkworms were reared on an artificial diet at temperatures of 24°C-27°C and 70%-90% humidity. Samples of total RNA were prepared from acutely dissected tissues from a silkworm, larval fat body on each stage, and Bm5 cells using Trizol and quantified by measuring absorbance at 260 nm. Typically, 2 mg of total RNA was reverse transcribed using random primers (GE Healthcare, Piscataway, NJ, USA) and mMLV reverse transcriptase (Invitrogen). RT-PCR using the forward primer FOR (5′-G ACT GAC AAT GGC AGA CAA TTC-3′) and REV (5′-CCG TCT AAT TAA TCG ACT CCT CAG-3′) for BmIHoP fragment was performed for 30 cycles (94°C for 30 s; 58°C for 30 s; and 72°C for 1 min (only 10 min in the final cycle)) with Taq DNA polymerase. A single band of BmIHoP amplicon (382 bp) was detected in RT-PCR.

Mammalian expression vector construction

The purified PCR product encoding BmIHoP was ligated at multiple cloning site of pCMV-Taq 2B vector (Addgene, Inc., Cambridge, MA, USA) with BamHI and Hind III. The accuracy of the ORF was further verified by sequencing. Transfections were performed with Fugene HD transfection reagent (Promega) with 2 μg of reporter plasmids per Ins1 cell-culture dish according to the manufacturer’s instructions.

Western blotting

Immunoblotting was performed according to the standard procedure secreted insulin from the Ins1 cell transfected by pCMV-Taq 2B-BmIHoP was collected and lysed by the addition of SDS sample buffer (62.5 mM Tris-HCl pH 6.8, 6% (w/v) SDS, 30% glycerol, 125 mM DTT, 0.03% (w/v) bromphenol blue) and separated by SDS-PAGE. The proteins were transferred to a nitrocellulose membrane, and the membrane was respectively incubated with the rabbit anti-polyclonal insulin antibody (Santa Cruz Biotechnology, Inc., Dallas, TX, USA), overnight at 4°C. The blots were developed using an enhanced chemiluminescence Western blotting detection system kit (Amersham, Uppsala, Sweden).

Results and Discussion

Diabetes is one of the most common life-related metabolic diseases, and its incidence is gradually increasing by every year. Korea has also experienced a gradual increase in the incidence rate of diabetes from an estimated less than 1% in 1970 to an estimated 8%-10% in 2000 [19]. Chronic serious complications in the diabetic patients occur as diseases of retina, kidney, peripheral neuron, and peripheral blood vessels [20]. Although there exist various chemicals for improving insulin resistance clinically, such long-term treatments are associated with side effects such as hypoglycemia, obesity, and other diabetes complications [21]. Henceforth, there exists a need for the development of novel anti-diabetic drugs from the natural substances including plants, organisms, and insects. Since ages, the silkworm, B. mori, has been used in the form of.
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In the present study, we concentrated on the islet homeostasis protein (IHoP), which is firstly reported as a novel factor that regulates glucagon synthesis in the pancreatic alpha-cells. Moreover, we isolated a cDNA encoding B. mori IHoP (BmIHoP, GenBank Accession No. KX893487) and described molecular characterization of BmIHoP and its differential gene expression in different tissues and at different developmental stages. A complete cDNA of BmIHoP shows a typical polyadenylation (AATAAA) signal at the end of 3′-terminal region and an open reading frame (ORF) of a putative protein consisting of 693 amino acids with a theoretical isoelectric point of 5.15 and a molecular weight of 77.5 kDa estimated by ExPASy. As shown in Figure 1, amino acid sequence of BmIHoP was analyzed based on reported homologues using Basic Local Alignment Search Tool (BLAST) of the NCBI. The amino acid residues of BmIHoP showed 52% identity with Homo sapiens, 52% with Mus musculus, 51% with Danio rerio, and 73% with Drosophila melanogaster. The result indicated that BmIHoP had been relatively well preserved beyond the species with at least 50% similarity, suggesting that despite little research on its biological function, BmIHoP is well demonstrated to play a vital role in the pancreatic alpha-cells.

The phylogenetic tree was generated based on amino acid substitution using to date reported IHoP homologues. The overall tree topology showed that BmIHoP of Lepidoptera (B. mori) is evolutionarily more closely related to the mouse (M. musculus) IHoP homologue than Diptera (D. melanogaster) IHoP homologue. It is interesting to note that BmIHoP shows a sister relationship with those of M. musculus IHoP homologues; however, rat (R. norvegicus) IHoP homologue shows evolutionary distance with BmIHoP (Figure 2). The information about their topology can greatly reduce experimental difficulties and lead to significant findings concerning their biological functions. Using ExPASy program, we have characterized BmIHoP on cellular localization and structural topology based on the amino acid sequences. Protein, BmIHoP, is located in the cytosol (estimated about 69%) and not in the transmembrane.
same level) without undergoing any drastic changes during developmental stages (Figure 3B). On the other hand, overall, no dramatic changes in BmIHoP gene expression in the instar were detected; however, relatively high expression of about 1.5-folds in the 5th-3rd instar was observed (Figure 3C).

Although the details of the molecular mechanism of BmIHoP are not currently understood, the result is that BmIHoP gene expression shows sever in the various tissues rather than at development stages.

OH et al. reported that IHoP is expressed in the pancreatic islet α-cells synthesizing glucagon, and it controls the cellular proliferation of β-cells under high glucose conditions [12]. It is suggested that IHoP could be a new functional protein for the maintenance of pancreatic homeostasis, and one of the new targets for type 1 diabetes treatment. We have constructed a cell line using Ins1 synthesizing insulin in which BmIHoP gene is over-expressed (BmIHoPover) to understand the association between BmIHoP and insulin secretion. It was observed that BmIHoP expression is greatly reduced (approximately half) insulin secretion from the β-cells (Figure 4A). Recently, it has been demonstrated that Sox17 (SRY-Box 17) is a key transcriptional regulator of the insulin trafficking and responsible for secretion of a hormone in the normal and pathologic mouse β-cells [18]. In this study, our data suggests that Sox17 expression is indirectly associated with the functional BmIHoP (Figure 4B) for the insulin secretion in the β-cells. A gene family of Bcl-2 encoding protein that can either suppress (e.g. Bcl-2 and Bcl-XC) or promote (e.g. Bid and Bak) apoptosis; It is also know that intracellular p53 protein promote apoptosis. The β-cells overexpressing BmIHoP induces only Bid expression in the MOCK, neither p53 nor Bcl-2. Eventually, the result can say that BmIHoP functionally induces apoptosis in the β-cells (Figure 4C).

In this study, IHoP homologue (BmIHoP) was isolated from B. mori; its gene expression was detected widely in all tissues revealing that the expression was regulated specifically, but not developmentally. Phylogenetic analysis revealed that BmIHoP shows a sister relationship with M. musculus IHoP homologue.

Figure 3. The expression level of BmIHoP mRNA in various tissues and developmental stages. (A) Various tissues were obtained skin, fat body, hemolymph, mid gut, post-silk gland, Malpighian, ovary and testis, respectively; (B) Samples were obtained at different developmental stages ranging from the stage of an egg to adult; (C) Samples obtained from 1st to 5th instars. The expression of 28S rRNA was used as an internal control for assessing the expression level of BmIHoP. RT-PCR was 3 times measured and those average values are represented by bar-graphs.

Figure 4. Forced BmIHoP expression in the β-cells. (A) Insulin secretion in the BmIHoP overexpressed β-cell by Western blotting; (B) Both the expression of SOX17; (C) Induction of apoptosis by RT-PCR. PPIA expression was used as an internal control for assessing the expression level of BmIHoP. Bar graphs represent the average of three RT-PCR values.

Overexpression of BmIHoP in the β-cells reduced insulin secretion and induced apoptosis. Overall, it is hypothesized that the present data on BmIHoP will provide more information on understanding pancreatic homeostasis. Eventually, it is
proposed that *BmIHoP* can potentially be applied for treatment of diet-induced obesity in the future.

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**References**


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