

Novel monoclonal antibodies against Pdx1 reveal feedback regulation of Pdx1 protein levels

T. Galbo,^{1,4} I.L. Pedersen,^{2,4}

T. Fløyel,³ C.H. Bang-Berthelsen,³ P. Serup,^{1,4} O.D. Madsen,^{1,4} J. Hald^{1,4}

¹Hagedorn Research Institute, Department of Beta-cell Regeneration, Gentofte, Denmark

²Novo Nordisk A/S, Department of Antibody Technology, Måløv, Denmark ³Hagedorn Research Institute, Department of Diabetes Genome Biology, Gentofte, Denmark

⁴Antibody Core, Beta Cell Biology Consortium, www.betacell.org

Abstract

The aim of this study was to characterize two monoclonal antibodies (F6A11 and F109-D12) generated against Pdx1 (pancreatic and duodenal homeobox-1), a homeodomain transcription factor, which is critical for pancreas formation as well as for normal pancreatic beta cell function. For production of monoclonal antibodies, we immunized Robertsonian POSF (RBF)mice with a GST-Pdx1 fusion protein containing a 68amino acid C-terminal fragment of rat Pdx1. These monoclonal antibodies detect Pdx1 by western blotting and allow immunohistochemical detection of Pdx1 in both mouse and rat tissue. F6A11 and F109-D12 produce IHC staining patterns indistinguishable from that obtained with highly specific polyclonal Pdx1 antisera raised in rabbits and goats, when applied to embryonic or adult mouse pancreatic tissue. In contrast to previously generated polyclonal anti-Pdx1 antisera, we also demonstrate that F6A11 works for intracellular fluorescence activated cell sorting (FACS) staining of Pdx1. By using F6A11, we characterize the induction of Pdx1 in the Doxycycline (DOX) inducible insulinoma cell line INSr $\alpha\beta$ -Pdx1 and follow the reduction of Pdx1 after removing Dox. Finally, we show that induction of exogenous Pdx1 leads to a reduction in endogenous Pdx1 levels, which suggests that a negative feedback loop is involved in maintaining correct levels of Pdx1 in the cell.

Introduction

The pancreatic duodenal homeobox gene 1, Pdx1, encodes a key regulatory factor required for pancreatic development in mouse^{1,2} and man,³ as well as for normal β -cell function including glucose-regulated insulin production.^{4,5} In the adult pancreas, high levels of Pdx1 expression are primarily restricted to insulin producing β -cells, and the protein was first identified and characterized as an insulin gene trans-activating factor.^{6,7} In the mature β -cell, Pdx1 is involved in glucose regulated insulin gene transcription⁸ and mutations in PDX1 cause maturity onset diabetes of the young (MODY).⁹

Pdx1 expression begins at embryonic day 8.5 (e8.5) in two ventro-lateral domains, before closure of the foregut endoderm. Slightly later (~e8.75) a dorsal expression domain appears at the level of somites 4 to 7. Until around e9.5, Pdx1 expression is restricted to the pancreatic buds, but from around e10.5 the Pdx1-expression domain expands to include the duodenum, the pyloric region and the antral stomach.

From e13.5-15.5, Pdx1 expression levels decline in the forming acinar cells, while remaining more broadly expressed throughout the developing ductal tree, where high-level Pdx1- and Nkx6.1-expression marks the forming β -cells.^{27,10,11} As the islets begin to form around e18.5, expression of Pdx1 becomes localized primarily to the β -cells.

Selective inactivation of Pdx1 expression in β -cells leads to loss of these and to diabetes.^{4,12} Furthermore, timed inactivation of Pdx1 expression during mouse pancreas development demonstrates Pdx1-requirement also for normal acinar development and maintenance of the exocrine transcription factor Ptf1a.⁵ Additionally, several target genes have been identified for Pdx1, including the insulin gene,⁸ *Nkx6.1*,¹³ *GLUT-2*,¹⁴ *IAPP*,^{15,16} glucokinase^{14,16} as well as somatostatin,^{17,18} exocrine genes such as Elastase1¹⁹ and finally also Pdx1 itself.^{20,21}

The rat α/β -cell hybrid subclone INSr $\alpha\beta$ -Pdx1 has been generated with the aim to further study Pdx1 regulated genes and the role of Pdx1 in β-cell maturation. In this cell line Pdx1 levels are under the control of the reverse tetracycline-dependent transactivator (rtTA).22 This INSr $\alpha\beta$ -Pdx1 cell line has been used to identify genes with potential functional significance in type 1 diabetes (T1D),23,24 to study insulin processing and secretion25,26 and to elucidate the role of Pdx-1 in β -cell maturation and acquired IL-1β-mediated toxicity.24 However, the details of the dynamic induction of Pdx1 in this important model system are not known, as well as the effects of expressing high levels of exogenous Pdx1 protein on the endogenous Pdx1 protein levels. By help of a more detailed biological

Correspondence: Jacob Hald, Hagedorn Research Institute, Niels Steensensvej 6, DK-2820 Gentofte, Denmark. Tel: +45.4443.9185 - Fax: +45.4443.8000 E-mail: hald@hagedorn.dk

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understanding of the system, more conclusive data may be drawn from such experiments. We used newly generated monoclonal antibodies against Pdx1 to study the induction of Pdx1 in the INSr $\alpha\beta$ -Pdx1 cell line and show that induction of exogenous Pdx1 leads to a reduction in endogenous Pdx1 levels. This suggests that a negative feedback loop is involved in maintaining correct levels of Pdx1 in the cell.

Materials and Methods

Expression and purification of GST-Pdx1

For immunizations, we cloned a 68 amino acid C-terminal fragment of the rat Pdx1 transcription factor (aa215-283) as a GST fusion protein.²⁷ The C termini of the rat, mouse and human Pdx1 proteins vary in sequence; the chosen GST-Pdx1 fusion protein have previously





allowed us to generate polyclonal antisera that may be either rat-specific or cross-react to mouse and/or human Pdx1.^{15,28,29} The GST-Pdx1 fusion protein was purified using a Glutathione Sepharose 4B Redipack column (Amersham Biosciences, Uppsala, Sweden).

Immunization

In order to obtain monoclonal antibodies, four RBF-mice³⁰ were immunized *i.p.* with 20 µg GST-Pdx1 biweekly. The first immunization was performed with complete Freund's adjuvant; this was followed by two injections in incomplete Freund's adjuvant. Ten days after the third immunization, the animals were eye-bled and the antibody response measured by ELISA. In addition, specific staining of β -cells of adult mice pancreatic tissue was determined by immunohistochemical staining. All animal experiments were performed according to national guidelines and approved by the national ethics committee.

Cell fusion

The mouse selected for generation of monoclonal antibodies was boosted i.v. with 10 μ g antigen in saline. Three days later, the spleen was harvested and used for hybridoma production. The 1.3×10^8 spleen cells were fused to 3.9×10^7 FOX-NY myeloma cells³⁰ using polyethyleneglycol (PEG) 1500 (Roche, Hvidovre, Denmark) as described in a previous study.³¹ The fused cells were initially seeded in 96 well tissue culture plates and the individual clones tested for the production of anti-Pdx1 antibodies, by ELISA and immunohistochemical staining. Positive clones were subcloned by limiting dilution.

ELISA

ELISA plates were coated O/N with 1 µg/mL GST-Pdx1 or pure GST (Sigma, Brøndby, Denmark) in PBS. Washing and blocking was performed with PBS, 0.05% Tween 20. Then, samples were added, after dilution in PBS, 0.5% Tween 20, 1 M NaCl, pH 7.2 and the plates incubated for 1 hr at room temperature (RT). The presence of antigen specific antibodies was detected with horse radish peroxidase (HRP) conjugated goat anti-mouse IgG (Caltag, Hamburg, Germany) diluted 1:2000. One-hundred µL of 3,3',5,5'-tetramethylbenzidine (TMB) (Kem-En-Tec, Copenhagen, Denmark) was added to each well and the reaction was terminated with 100 µL 4 M H₃PO₄. The absorbance of the ELISA plates was read at 450 and 620 nm.

Isotyping and antibody purification

The isotypes of the antibodies were determined using the Mouse Monoclonal Antibody Isotyping Kit (Boehringer Mannheim, Hvidovre, Denmark), according to the manufacturer's instructions. The antibodies were purified from culture supernatant using a protein A Sepharose 4 Fast Flow column (Amersham Biosciences, Uppsala, Sweden). The pH of the filtered supernatants was adjusted to 8.5 and the solutions supplemented with 3 M NaCl. The samples were applied to the column, initially equilibrated and subsequently washed with 50 mM Tris, 3.5 M NaCl, pH 8.5. Elution was performed with 25 mM Citric acid, 4.5 mM Trinatrium-citrate and pH 3. The pooled eluates were dialyzed against phosphate buffered saline (PBS) and 0.05% NaN3 was added to the purified antibodies.

Immunohistochemistry

In general, animal sera, hybridoma supernatants and purified antibodies against Pdx1 were analysed by immunohistochemical staining on frozen sections of adult murine pancreas. Tissue was fixed in 4% PFA O/N, cryo-protected by O/N incubation in 30% sucrose and finally embedded in Tissue-Tek. Ten µm sections were cut and stored at -80°C. For antigen retrieval, the sections were soaked in 200 mL 0.01M Citrate buffer and microwaved for 4 min at 600W, followed by 15 min at 250W and finally left cooling for 20 min. The tissue sections were rinsed in PBS, treated with 3% H₂O₂ for 5 min and rinsed again. TNB blocking buffer was added and the sections blocked for 30 min, before O/N incubation with the mouse anti-Pdx1 antibodies at a concentration of 0.5 µg/mL, diluted 1:5000 in TNB blocking buffer. For absorption studies, the anti-Pdx1 antibodies were pre-incubated with GST-Pdx1 or the GST-Nkx6.1 before use, as described by previous studies.^{28,32} The sections were washed 3×5 min in PBS between the incubations with primary antibody and the following incubations with biotinylated secondary anti mouse antibody (Zymed) for 30 minutes and streptavidin-peroxidase conjugate (Zymed) for 15 minutes. The bound antibodies were visualized with the Cv3 fluorophore tyramide signal amplification (TSA) agent (PerkinElmer) diluted 1:100, after a 10 min incubation period and a subsequent wash in PBS. Double staining with antisera raised against Pdx1 in different species was performed with relevant mouse anti-Pdx1 clone and rabbit anti-Pdx1 1858.5, 1:800.29 Triple staining experiments were performed by first adding Pdx1 antibody and developing it with Cy3- TSA, then incubating with guineapig-antiinsulin 1:100 (Abcam, ab7842) and rabbit anti glucagon 1:300 (DAKO, A0565), visualized by donkey anti guineapig-Cy2 and donkey anti rabbit-Cy5 (Jackson Immunoresearch). Whole mount stains were done as described by Ahnfelt-Ronne J et al.33 Mouse anti-Pdx1 (F6) 1:700, guineapig anti-glucagon 1:4000, goat anti-Pdx1 1:15000 (kind gift from Chris Wright)

and developed with donkey anti rabbit-Cy3, donkey anti-guinea pig-Cy2 and donkey anti-goat-Cy5. Images were collected with on a Zeiss LSM 510 META confocal microscope.

Nuclear extracts

The SV40 large T-antigen transformed BTC6 insulinoma cells³⁴ and aTC1.9³⁵ were grown in DMEM (glucose 1g/L) supplemented with 1% glutamax 5% FCS, 20 mM Hepes, 10 µM 2-mercaptoethanol and 1% glutamax, 10 % FCS, respectively. Nuclear extracts from the cells were generated from cells harvested by trypsination. Cell pellets were lysed in 1 mL lysis buffer (10 mM Hepes pH 7.9, 10 mM KCL, 0.1 mM EDTA and 0.1% Nonidet P40 [NP40], supplemented with the protease inhibitor cocktail containing 1 mM AEBSF, 1 mM DTT, 10 ug/mL Aprotinin, 10 µg/mL Leupeptin and 1 µg/mL Pepstatin A). The cell extract was vortexed and incubated on ice for 45 min. After centrifugation, the pellet was re-suspended in 20 mM Hepes pH 7.9, 0.4M NaCl and 1 mM EDTA supplemented with protease inhibitor cocktail. After five min incubation on ice, the extract was centrifuged and the protein concentration of the supernatant was measured by the Bradford method (Bio-Rad Laboratories, Herlev Denmark).

Western blots

Purified GST-Pdx1 fusion protein or 10 µg of nuclear extract was heated for 10 min at 70°C in sample buffer and loaded onto NuPAGE SDS PAGE gels, run alongside the MagicMark XP Standard (Invitrogen, Taastrup, Denmark). The proteins were blotted onto a nitrocellulose membrane using the NuPAGE 4-morpholinepropanesulfonic acid (MOPS) buffer system. The membrane was washed in PBS 0.1%Tween and blocked for 1 hr at RT in Amersham Membrane blocking agent NIF833. 5% w/v in PBS 0.1% Tween. The monoclonal anti-Pdx1 antibodies were diluted to a concentration of 1 ug/mL before O/N incubation with the membrane at 4°C. A HRP-conjugated rabbit Anti-Mouse IgG (DakoCytomation, Glostrup, Denmark) was used as the secondary antibody at a dilution of 1:5000, for 1 hr at RT. Chemiluminescence was detected using the ECL analysis system (Amersham Biosciences, Uppsala, Sweden) and developed with a LAS 3000 from FUJI FILM.

Pdx1 induction

INSrαβ-Pdx1²² cells, passage 38-42, were stimulated for 24 hours in medium (RPMI 1640 supplied with 10% Fetal Bovine Serum (Gibco), 1% Penicillin/Streptomycin (Gibco) and 50 μ M β-mercaptoethanol (Sigma) with 500 ng/mL doxycycline (Dox) (Sigma). Cells were rinsed for 5 min in PBS w/o Ca²⁺/Mg²⁺ and for one hour



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in fresh medium without Dox. The cells were then given fresh medium without Dox. Cells were harvested 0, 4, 24, 28, 32, 48, 60 and 72 hours after stimulation with 0.05% Trypsin-EDTA (Gibco) and fixated for 20 minutes in 4% paraformaldehyde (Lilly's fixative) (VWR International).

Pdx1 stainings for FACS analysis

For each time point, 10⁶ fixed cells were permeabilised by 0.03% Triton X-100 (Sigma) in PBS w/o Ca2+/Mg2+ supplemented with 0.1% Bovine Serum Albumin (BSA) (Sigma) for one hour. In order to block unspecific binding, 10% donkey serum (Jackson) was added for additionally one hour. Cells were then incubated overnight with mouse monoclonal anti-Pdx1 antibody (F6A11) diluted to a final concentration of 5.00 µg/mL, or mouse IgG1k (Abcam) diluted to a final concentration of 5.00 µg/mL. Rabbit anti-Pdx1 serum (1858.5) or nonimmune rabbit serum were tested at dilutions of 1:1000 and 1:5000. The cells were washed three times in PBS + 0.1% BSA, then incubated with a Cy2 conjugated donkey anti-mouse antibody or a Cy2 conjugated donkey anti-rabbit antibody (both Jackson ImmunoResearch) diluted 1:300, for one hour. Cells were then washed three times in PBS + 0.1% BSA and analyzed on a FACS Aria flow cytometer (Beckton-Dickinson). Data analysis was performed using FCS Express Version 3 (De Novo Software).

Real-time qPCR

After stimulation with Dox, total RNA was extracted from the INSr $\alpha\beta$ -Pdx1 cells and cDNA was prepared using reverse transcription, as described by the manufacturer (TagMan RT reagents, Applied Biosystems, Foster City, CA, USA). The expression levels of the genes were analysed by use of TaqMan assays (Assay ID: Rat Pdx1 Rn00755591 m1, Mouse Pdx1 Mm00435565_m1, Rat Ppia Rn00690933_m1, Applied Biosystems) and samples were run on a TaqMan 7900HT system (Applied Biosystems). The expression level of the gene of interest was normalised against the expression of the housekeeping gene peptidylprolyl isomerase A (Ppia) and evaluated using the delta-delta Ct method.36

Image capturing and handeling

All immunofluorescence images were recorded on a LSM510 confocal microscope (Carl Zeiss; Brock & Michelsen, Birkerød, Denmark). Image manipulations were performed using linear adjustments over the entire image in LSM Zeiss viewer and Adobe Photoshop.



Figure 1. GST-Pdx1 is of expected size and the monoclonal antibodies F6 and F109 recognize Pdx1 from β TC6 with correct size; furthermore, F6 and F109 specifically recognize Pdx1 in adult mouse pancreas. (A) SDS PAGE analysis of the purified GST-Pdx1 fusion protein, demonstrating the size and purity of the antigen. For the non-reduced protein a double band around 37 kDa is observed (lane 1), whereas under reducing conditions only a single band around 38 kDa is seen (lane 2). In addition, a minor band of approximately 70 kDa is present, which may represent a dimer (filled-in arrowhead). Western blotting of the GST-Pdx1 using the monoclonal anti-Pdx1 antibody F6 revealed that both bands seen on the SDS PAGE gel were derived from the Pdx1 fusion protein (lane 3). The hollow arrow head point to the 70 kDa that is also recognized by F6. (B) Western blot analysis of the purified Pdx1 antibodies F109 (lane 1 and 2) and F6 (lane 3 and 4) for binding to nuclear extracts from β TC6 cells (lane 2 and 4) and α TC1.9 cells (lane 1 and 3) at 10 ug/well, demonstrating that the anti-Pdx1 antibodies identify a single band around 39 kDa corresponding to the Pdx1 protein. This binding could be absorbed with the GST-Pdx1 protein, but not with an irrelevant GST fusion protein (data not shown). (C, D) Immunohistochemical staining of frozen adult mouse pancreas with the purified monoclonal antibodies F6 (C, E and G) and F109 (D, F and H) result in nuclear reactivity in the endocrine islets of Langerhans (C and D). This staining was absorbed by preincubation of the antibodies with the GST-Pdx1 fusion protein - islet is outlined by broken line (E and F), but not by an irrelevant GST fusion protein (G and H). The magnification of histochemical stainings is the same. Scale bar is 50 µm.



Results

A detailed understanding of the dynamic changes in Pdx1 levels in the INSr $\alpha\beta$ -Pdx1 cell line after induction of Pdx1 is vital to data interpretation from this system. To characterize it, we induced INSr $\alpha\beta$ -Pdx1 with Dox for 24 hours, washed the Dox out and followed the cells for further 48 hours. To obtain a quantitative measure of Pdx1 levels in the INSrαβ-Pdx1 cell line, we assayed Pdx1 levels by FACS on permeabilized cells. Using the validated 1858.5 rabbit anti-Pdx1 antiserum,29 we were unable to detect a shift in fluorescence after 24 hours of Dox induction. However, using the 1858.5 rabbit anti-Pdx1 antiserum on INSraß-Pdx1 cells in immunocytochemistry (ICC) did result in clear nuclear Pdx1 staining (data not shown). This is likely due to the use of antiserum instead of the monoclonal antibodies that are usually utilized in FACS experiments with success. However, since no good monoclonal antiPdx1 antibody was available, we set out to generate one.

Monoclonal antibodies against the Pdx1 transcription factor were generated by immunising RBF mice with a GST-Pdx1 fusion protein, containing the 68 amino acids of the C-terminal fragment of rat Pdx1 protein (aa 215-283). We have previously used this fusion-construct²⁷ to generate a series of high-titered rabbit antisera, some of which showed cross-reaction to both mouse and human Pdx1.^{15,28,29} The expected size of the GST-Pdx1 fusion protein is ~37 kDa. The purified protein is observed as a double band around 37 kDa (Figure 1A, lane 1), which however could be reduced to a single band of approximately the expected molecular weight (Figure 1A, lane 2). In addition, a band of ~70 kDa was observed (Figure 1A, lane 1 filled-in arrow head). This larger band may represent a dimer, as another anti-Pdx1 antibody, when analyzed by western blotting (Figure 1A, lane 3 hollow arrow head), also recognized it.

Three out of four mice responded to immunization with antibody titers higher than 1:36.450, as determined by binding to the GST-Pdx1 fusion protein in direct ELISA. However, only one of the sera stained a sub-population of cells on sections of adult mouse pancreas at a dilution of 1:100, corresponding to the β -cell population. Thus, this mouse was selected as the best starting material for hybridoma generation, to develop monoclonal antibodies against Pdx1 that could be used primarily for FACS but also for other immuno-techniques.

The resulting hybridomas were screened for the selective binding to Pdx1, by binding to the rat GST-Pdx1 fusion protein vs GST in direct ELISA. Subsequently, the 74 most positive clones were further screened for staining of PFA fixed, frozen sections of adult mouse pancreas, by immunohistochemstry. Five hybridomas pro-



Figure 2. Monoclonal immunoreactivity co-localize with immunoreactivity generated by validated anti-Pdx1 sera on sections and in whole mount IHC, and co-localize with β -cells, but not α -cells. Staining of adult pancreas with the monoclonal anti-Pdx1 antibodies F6 (B) and F109 (E) showed a high degree of overlap with the previously described 1858.5 anti-Pdx1 antiserum (A and D), as seen in C and F (double exposures). Whole-mount triple staining of the e10.5 budding stage of pancreas development (G-I) by confocal microscopy: Polyclonal goat anti-Pdx1 (green channel, G); F6 monoclonal anti-Pdx1 (red channel, H); guinea-pig anti glucagon (blue channel in I). Again almost complete immunoreactive overlap in the pancreatic domain is observed between the monoclonal F6 and the goat anti-Pdx1 antiserum (H vs G) – also visualized as yellow in overlay (I). It is also evident that all glucagon positive cells (I, blue cells) are devoid of Pdx1 immunoreactivity. Double- (J, K) or triple-staining (L) of frozen adult mouse pancreas with the monoclonal anti-Pdx1 antibodies F6 (J and L) and F109 (K) with anti-insulin antibody (J, K, L green) and anti-glucagon antibody (L, blue). The monoclonal anti-Pdx1 antibodies recognize islet nuclei in the mature islet β -cells expressing insulin in the cytosol, but not the glucagon producing α -cell. Scale bars: 50 μ m except panel l, which is 25 µm.

duced very strong signals, whereas further four gave weaker signals. Additional subcloning of the cells resulted in two stable hybridomas producing high affinity mouse monoclonal antibodies against Pdx1, termed F6A11 and F109-D12 (hereafter: F6 and F109). These two hybridomas were produced in large scale and the antibodies purified from the cell culture supernatant using protein A. The antibodies produced by both the F6 and F109 hybridomas were determined to be of the IgG1/ κ subtype (*data not shown*).

Both the monoclonal anti-Pdx1 antibodies F6 and F109 were shown to recognize the GST-Pdx1 antigen by western blotting (Figure 1A, lane 3, and data not shown). In addition, the antibodies stained a band of 39 kDa, correspon-



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ding to the Pdx1 protein, in nuclear extracts from the SV40 large T-antigen transformed mouse insulinoma β TC6 cell line (Figure 1B, lane 2 and 4), but not from the α TC1.9 gluca gonoma mouse cell line (Figure 1B, lane 1 and 3). This again reflects the expected expression of Pdx1 in these two islet tumor lines. This Western-blot staining specificity of F6 and F109 was confirmed by pre-absorption studies, where only pre-incubation with GST-Pdx1 fusion protein and not GST-Nkx6.1 was able to abolish the appearance of the 39 kDa band (*data not shown*).

The purified monoclonal mouse anti-Pdx1 antibodies F6 and F109 were further characterized by staining of frozen sections of adult mouse pancreas and fetal pancreas. Both antibodies were demonstrated to specifically stain the nuclei of a sub-population of cells in the endocrine islets of Langerhans, as expected for a Pdx1 reactive antibody (Figure 1, C and D). Specificity was demonstrated as the staining of the cell nuclei was eliminated by pre-absorption of the antibodies with the GST-Pdx1 fusion protein (Figure 1, E and F), but not by pre-absorption with an irrelevant GST fusion protein (Figure 1, G and H). Furthermore, double staining of adult pancreas with a rabbit anti-Pdx1 antiserum (1858.5)²⁹ demonstrated a near complete overlap of the nuclei recognized by the two antibodies (Figure 2A-C and D-F). On close inspection of the co-stain (Figure 2 A-F) and other data (not shown), we found that the F6 monoclonal antibody was superior regarding: i) less background and more consistent performance in IHC and ii) an apparent better growthrate of the hybridoma. We thus went on to further characterize this antibody and performed triple staining of mouse e10.5 tissue with F6, goat anti-Pdx1 and guinea-pig anti-glucagon. This again confirmed that F6 and goat anti-Pdx1 stain in overlapping patterns (Figure 2G-I). However, the goat serum appears to give a stronger signal in the duodenum and stomach (Figure 2G-I). Co-staining of adult mouse pancreas for Pdx1, insulin and glucagon, clearly demonstrated the expected β -cell specific staining pattern, as insulin positive cells are positive for Pdx1 immunoreactivity (Figure 2J-K), while glucagon positive cells are not (blue) (Figure 2L). The antibodies were also shown to react with rat Pdx1, but they did not recognize Pdx1 from human tissue (data not shown). Furthermore, the antibodies worked equally well on frozen/fixed or paraffin sections as well as by using Lilly's fixative (data not shown). Finally, the antibodies were efficient when used in a standard indirect immunofluorescence experiments; i.e. without the use of TSA amplification (data not shown).

Having generated and validated monoclonal antibodies to Pdx1, we then wanted to test if they were useful for assaying Pdx1 protein lev-



Figure 3. Dynamic induction and negative feedback regulation of Pdx1 levels in INSr $\alpha\beta$ -Pdx1. The induction of Pdx1 in INSr $\alpha\beta$ -Pdx1 cells is followed by FACS analysis using the monoclonal antibody F6 (A, B, and C). A rapid induction of Pdx1 by Dox can be seen with a clear increase in fluorescence intensity from 0 hours (red line) to 4 hours (blue line) before the Pdx1 level peaks after 24 hours (purple line) (A). Following removal of Dox, the Pdx1 level slowly begins to decline, which can be seen after 28 hours (red line) (4 hours after removal of Dox) (B). The Pdx1 level continues to decline from 28 hours (red line) to 32 hours (blue line) (8 hours after removal of Dox), before eventually reaching the lowest point after 48 hours (purple line) (24 hours after removal of Dox) (B). Slowly, from 48 hours (purple line (B)) to 60 hours (red line (C)) (36 hours after removal of Dox) and through to 72 hours (blue line (C)) (48 hours after removal of Dox), the Pdx1 level recovers and gradually approaches the initial level found at 0 hours (red line (a)). Panel a-c constitutes a single experiment, representative of three independent experiments. The bars in figure G represent averages of the relative Pdx1 levels found in the three independent experiments. Student's T-test assuming unequal variances was applied to test whether Pdx1 levels at the time points were different to the level at 0 hours"*" or 24 hours (time of Dox removal)"#". *P<0.05, **P<0.01, #P<0.05, ##P<0.01, ###P<0.005. Confocal microscopical analysis of the samples analysed on the FACS and later supplied with DNA binding fluorophore DAPI confirm the FACS data (D, E and F). The images reveal a visible increase in Pdx1 from 0 hours (D) to the peak at 24 hours (E) and clear decline to 48 hours (F). Error bars are sd.



els in individual cells by intracellular FACS analysis. To this end, $INSr\alpha\beta$ -Pdx1 cells were induced with Dox for 24 h and Pdx1 levels were analysed at 0 h. 4 h and 24 h by FACS. The Dox was then washed out and, the cells were followed for additionally 48 h and tested in FACS at 4, 8, 24, 36 and 48 h post-wash. A fast and significant rise in mean fluorescence intensity was observed already after 4 h, reaching a 2.3fold increase compared with control levels at 24 h (Figure 3A and G). Increase in Pdx1 levels was confirmed by western blotting (data not shown). Pdx1 levels declined significantly already 8 h after removal of Dox and a further reduction was observed 24 h after Dox removal. At this time, the Pdx1 level had been reduced 3.4-fold compared with the maximum levels 24 h after addition of Dox. Surprisingly, Pdx1 levels were reduced significantly below starting levels 24 h after removal of Dox. However, the reduction in Pdx1 levels observed at 24 h after Dox removal was normalized after additional 24 h (Figure 3 G). In order to confirm that the fluorescence-signal observed by FACS reflected an immunoreaction to Pdx1, we investigated the staining in the microscope and found a convincing nuclear staining at 24 h after induction with Dox (Figure 3E). Furthermore, the strong induction of Pdx1 was confirmed by comparing the immunoreaction before and after induction (Figure 3 D vs. E). The observed decline in Pdx1 levels at 24 h was confirmed by fluorescence microscopy when comparing this time point with cells before induction (Figure 3D vs. F).

To investigate whether the observed reduction in Pdx1 levels 24 h after Dox removal (Figure 3F and G) might reflect a reduction in the synthesis of endogenous rat Pdx1, we performed qPCR for mouse (induced) and rat (endogenous) Pdx1 mRNA at 24 h after induction, the time point where total Pdx1 protein level peaks. This analysis showed that mouse Pdx1 mRNA was up-regulated 28-fold (Figure 4, black). In contrast, rat Pdx1 mRNA was reduced by 3 fold (Figure 4, grey). To rule out a general reduction of all endogenous genes upon induction with Dox, gPCR was performed for several other mRNAs. The other profiled genes did not show any significant differential regulation (data not shown).

Discussion

The two new Pdx1-specific monoclonal mouse antibodies, F6 and F109, could hence be added to our previously published panel of reagents for the immunohistochemical detection of critical pancreatic transcription factors including Ngn3,³⁷ Nkx6.1³² and Ptf1a.³⁸ These monoclonals are important tools for investigations aimed at understanding the mechanisms



Figure 4. Rat Pdx1 mRNA expression declines upon exogenous expression of mouse Pdx1. INSr $\alpha\beta$ -Pdx1 cells were cultured with/without Dox for 24 hours prior to RNA extraction. Mouse Pdx1 (black) and rat Pdx1 (grey) mRNA expressions were analysed by real-time qPCR and the data was normalised to the housekeeping gene Ppia. The results represent means±sem.

of pancreatic development from the earliest budding stages to the final differentiation of mature insulin secreting β -cells. In particular, they will be of great value in FACS studies and for multi-label immunohistochemistry. Polyclonal antisera against Pdx1 have been available for more than 10 years and to our knowledge F6 and F109 represent the first monoclonal anti-Pdx1 antibodies useful for FACS as well as for immunohistochemical applications.

In this study, we used the F6 monoclonal antibody to characterize the induction of Pdx1 protein in INSrαβ-Pdx1 cells by fluorescence-activated cell sorting (FACS) analysis. Our FACS results show that the fluorescence intensity before induction follows a normal distribution as expected for a homogenous cell population. After induction with Dox, the normal distribution of the fluorescence intensity is maintained as the mean fluorescence intensity increases and subsequently decreases, indicating that the entire INSr $\alpha\beta$ -Pdx1 cell population responds homogenously. Furthermore, our FACS results show that Pdx1 is induced within hours after addition of Dox and continues to increase up to 24 h. Removal of Dox results in a decrease in Pdx1 levels, which reach their lowest point 24 h after removal of Dox. Surprisingly, at this point we find Pdx1 levels to be significantly lower than before Dox stimulation (0 h), indicating a possible negative feedback loop. A substantial amount of existing literature shows that Pdx1 can bind to its own promoter sequence^{21,39,40} and a role for Pdx1 in controlling its own promoter activity through a positive feedback mechanism has previously been suggested.20,40 However, our finding that Pdx1 has a repressive effect upon its own expression is further supported by qPCR data that show a 3-fold decline in endogenous Pdx1 mRNA 24 h after induction with Dox. This 3-fold decline in endogenous Pdx1 mRNA could be due to general squelching resulting from the over expression of mouse Pdx1 by addition of Dox. However, an array study was performed on

the INSrab-Pdx1 cells to exclude this possibility. Twenty six hours after induction with Dox, 16% of the probe sets were differentially regulated and 48 hours after induction, only 12% of the probe sets were differentially regulated (H. Bang-Berthelsen, manuscript in preparation). Among the differentially regulated genes, there are many genes expected to be regulated negatively or positively by Pdx1, substantiating the validity of the system. Yet, further studies are needed to clarify the mechanism of this novel negative feedback loop. We propose a model in which increased amounts of Pdx1 protein lead to an up-regulation of a negative regulator of the Pdx1 gene, resulting in the normalization of Pdx1 protein levels.

The availability of these new monoclonal antibodies through the Developmental Studies Hybridoma Bank (http://dshb.biology. uiowa.edu) will aid such studies.

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