Insulin immuno-neutralization in chicken: effects on insulin signaling and gene expression in liver and muscle

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Abstract

In order to evaluate the role of insulin in chicken, an insulin immuno-neutralization was performed. Fed chickens received 1 or 3 i.v. injections of anti-insulin serum (2-h intervals), while fed or fasted controls received normal serum. Measurements included insulin signaling cascade (at 1 h in liver and muscle), metabolic or endocrine plasma parameters (at 1 and 5 h), and qRT-PCR analysis (at 5 h) of 23 genes involved in endocrine regulation, metabolisms, and transcription. Most plasma parameters and food intake were altered by insulin privation as early as 1 h and largely at 5 h. The initial steps of insulin signaling pathways including insulin receptor (IR), IR substrate-1 (IRS-1), and Src homology collagen and downstream elements: phosphatidylinositol 3-kinase (PI3K), Akt, GSK3, ERK2, and S6 ribosomal protein) were accordingly turned off in the liver. In the muscle, IR, IRS-1 tyrosine phosphorylation, and PI3K activity remained unchanged, whereas several subsequent steps were altered by insulin privation. In both tissues, AMPK was not altered. In the liver, insulin privation decreased Egr1, PPARγ, SREBP1, THRSPα (spot14), D2-deiodinase, glucokinase (GK), and fatty acid synthase (whereas D3-deiodinase and IGF-binding protein1 transcripts were up-regulated. Liver SREBP1 and GK and plasma IGFBP1 proteins were accordingly down- and up-regulated. In the muscle, PPARβ and atrogin-1 mRNA increased and Egr1 mRNA decreased. Changes in messengers were partly mimicked by fasting. Thus, insulin signaling in muscle is peculiar in chicken and is strictly dependent on insulin in fed status. The ‘diabetic’ status induced by insulin immuno-neutralization is accompanied by impairments of glucagon secretion, thyroid axis, and expression of several genes involved in regulatory pathways or metabolisms, evidencing pleiotropic effects of insulin in fed chicken.

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Introduction

Despite the presence of hyperactive endogenous insulin circulating at ‘normal’ concentrations, chickens mimic a mammalian type 2 diabetic status by showing high glycemia and low sensitivity to exogenous insulin (Simon 1989, Akiba et al. 1999, Tokushima et al. 2003). Insulin dependency of chickens in the control of glycemia has, however, been shown in the fed state following extensive pancreatectomy or insulin immuno-neutralization (Simon 1989, Simon et al. 2000). In the fasted state, the control of glycemia is in contrast dependent on glucagon since fasted pancreatectomized chickens tend to develop hypoglycemia rather than hyperglycemia (Simon 1989). Similar conclusions have been obtained in ducks (Mirskey et al. 1964, Simon 1989 for a review).

Insulin receptor (IR) structure and two IR substrates have been characterized in chicken tissues: IR substrate 1 (IRS-1) and Src homology collagen protein (Shc) (Simon & Taouis 1993, Taouis et al. 1996). Tyrosine phosphorylation of IR β-subunit, IRS-1 and Shc, and phosphatidylinositol 3-kinase (PI3K) activity appeared responsive to insulin status in the liver under several experimental conditions (fasting/re-feeding, chronic corticosterone treatment, genetically fat or lean chickens, or exogenous insulin challenge (Dupont et al. 1998a,b,c, 1999, 2004)). In contrast, these signaling steps (with the exception of Shc) were not altered in leg muscles by the various experimental conditions. This peculiar feature is consistent with an absence of change in tyrosine kinase activity observed for muscle IR purified from fasted/re-fed chickens (Adamo et al. 1987). However, in the chicken muscle, Shc
tyrosine phosphorylation is stimulated by re-feeding or exogenous insulin (Dupont et al. 1998b, 2004). In addition to this apparent insulin refractoriness of muscle PI3K activity mentioned earlier, another peculiarity comes from the amount of p85 subunit and the activity of PI3K, which are largely higher in chicken muscle, when compared with the rat under the same conditions (Dupont et al. 2004). The apparent independence between insulin status and the level of IRS-1 tyrosine phosphorylation has been recently confirmed in chicken pectoral muscle, a pure glycolytic-type muscle (Duchène et al. 2008a). To better understand the mechanisms of insulin action in chicken muscle, other components of insulin signaling have been recently characterized, namely the PKB/Akt, P70S6K1, and ERK2 kinases (Bigot et al. 2003a,b, Duchène et al. 2008a,c). In mammals, activations of PKB/Akt and P70S6K1 are considered to be under the control of PI3K (White & Kahn 1994). In chicken muscle (leg or pectoral muscle), PKB/Akt, P70S6K1, and ERK2 were all activated by re-feeding or exogenous insulin (Bigot et al. 2003a,b, Duchène et al. 2008a,c). Under these conditions, an increase in the phosphorylation of IRS-1 on Ser 632/635 was observed in chicken muscle (Duchène et al. 2008c). These specific serine residues are involved in the development of insulin resistance in mammals (Bouzakri et al. 2003). This unique feature of chicken muscle, an apparent refractoriness of early insulin signaling steps associated with normal sensitivity of more distal steps, represents a challenging model for insulin action. In the absence of an experimental chicken diabetes model (diabetogenic drugs are ineffective in this species as shown previously by Simon (1989)), we investigated the effect of insulin immuno-neutralization on insulin signaling cascade (including GSK3) and gene expression in the liver and the muscle in order to evaluate the action of insulin in fed chickens.

Materials and Methods

Experimental protocol

One-day-old male broiler chickens (n = 75, ISA 915, Institut de Sélection Animale, Saint Brieuc, France) were housed in an environment-controlled room in individual wire cages equipped with individual feeders and water bowls. They were fed a conventional balanced diet ad libitum and after 2 days of age, exposed to a 14 h-light period per day (0600–2000 h). Diet (3050 kCal or 12.8 mJ/kg, metabolizable energy, 22% proteins (N×6.25)) were based on corn, wheat, peas, soya bean meal, corn gluten, and rapeseed oil. Body weight was measured at 1, 8, 12, and 14 days to minimize handling stress and individualize five groups of seven chickens exhibiting similar body weights (BW) at 16 or 17 days of age. The fed control group received three i.v. injections delivering normal guinea pig serum (PromoCell, Heidelberg, Germany; 1.5 ml/kg each) at time 0, 2, and 4 h (abbreviated as 5hrfed-Ab group, BW = 475 ± 18 g, n = 7). Another fed group received three i.v. injections delivering anti-porcine insulin guinea pig serum (1.5 ml/kg) at 0, 2, and 4 h (abbreviated as 5hrfed-Ab group, BW = 500 ± 22 g, n = 7). Immune sera were prepared as described earlier (Simon et al. 2000). Two additional treatment groups, maintained on feed, received one single i.v. injection (1.5 ml/kg) at 4 h of either normal guinea pig serum (abbreviated as 1hrfed-C group, BW = 505 ± 21 g, n = 7) or anti-porcine insulin guinea pig serum (abbreviated as 1hrfed-Ab group, BW = 496 ± 17 g, n = 7). The last group, which served as an additional control to measure the extent of changes induced by insulin immuno-neutralization, was fasted for 5 h and given three i.v. injections of normal guinea pig serum (1.5 ml/kg) at time 0, 2, and 4 h (abbreviated as 5hrfasted-C group, BW = 471 ± 18 g, n = 7). In a preliminary assay, it was checked that the anti-insulin antibody mixture had a hyperglycemic effect (and not an insulin mimetic effect). Food intake was measured after 1 h for 1hrfed-C and 1hrfed-Ab groups or 5-h insulin immuno-neutralization for 5hrfed-C and 5hrfed-Ab groups. After blood sampling under EDTA, the chickens were killed by cervical dislocation. Liver and leg muscle samples were quickly removed, frozen in liquid nitrogen, and stored at −80°C until analysis. After blood centrifugation, aliquots of plasma were prepared and stored at −20°C until analysis. All procedures described herein were approved by the Agricultural Agency and the Scientific Research Agency and conducted in accordance with the guidelines for Care and Use of Agricultural Animals in Agricultural Research and Teaching.

Metabolic and hormone analyses

Plasma glucose levels were measured by the glucose oxidase method (Glucose Beckman Analyser 2, Beckman, Palo Alto, CA, USA). Free or non-esterified fatty acid (NEFA) levels were determined with an enzymatic colorimetric kit (Wako Chemicals, Neuss, Germany). Plasma αNH2NPN concentrations (α-amino-non-protein nitrogen, an estimate of total free amino acids) were measured after extraction with 10% (v/v) sulfosalicylic acid, using the 2% ninhydrin reagent (Sigma Chemicals) and i-serine as standard. Plasma glucagon levels were determined by RIA with a specific C-terminal pancreatic glucagon antibody, which does not cross-react with gut glucagon (Ruflié et al. 1998), a generous gift from Leclercq-Meyer, and porcine glucagon (Elli Lilly France) as the standard. Plasma tri-iodothyronine (T3) concentrations were estimated by specific RIA using a commercial coated tube kit ( Coat-A-Count, Diagnostic Product Corporation France, La Garenne Colombes, France). The RIA appeared not sensitive enough to get reliable plasma thyroxine (T4) concentrations using the present samples. For each hormone, all samples were measured in one RIA.

Preparation of tissue extracts

For analyzing insulin signaling components, the liver and leg muscle soluble protein lysates were prepared as described...
previously (Dupont et al. 1998a,b). Protein content of extracts was determined using the BCA kit (Interchim, Montluçon, France). For preparing liver membranes and nuclear extracts, liver samples were homogenized in buffer 1 (10 mM HEPES, pH 7.9, 10 mM KCl, 1.5 mM MgCl2, 1 mM EDTA, 1 mM EGTA, 1 mM dithiothreitol). Homogenates were centrifuged at 1100 g for 10 min. Resulting nuclear pellets were washed once in buffer 1 and then re-suspended in buffer 2 (20 mM HEPES, pH 7-9, 420 mM NaCl, 25% (v/v) glycerol, 1.5 mM MgCl2, 1 mM EDTA, 1 mM EGTA, 1 mM dithiothreitol). This suspension was rotated for 30 min and then centrifuged at 15 000 g for 30 min. The resulting supernatant was designated as the nuclear extract fraction. The membrane extract fraction was prepared by centrifuging the supernatant of the original 1100 g spin for 1 h at 100 000 g. The resulting membrane pellet was dissolved in buffer 3 (10 mM Tris, pH 6-8, 100 mM NaCl, 1% (w/v) SDS, 1 mM EDTA, 1 mM EGTA, 1 mM dithiothreitol).

Western blotting and PI3K activity

Tissue extracts were subjected to electrophoresis on 10% (w/v) SDS-polyacrylamide gels under reducing conditions and electro-transferred as described previously (Dupont et al. 1998a). Membranes were then incubated overnight at 4 °C with appropriate antibodies: p-ERK1/2 (T202/Y204), p-Akt (S473), Akt, p-GSK3-β (S9), p-P70S6K1 (T389), p-P70S6K1 (T421/S424), and p-S6 (S235/236) (Cell Signaling, Beverly, MA, USA); Shc and phosphotyrosine (PY20) (Biosciences, Le Pont de Claix, France); ERK2, P70S6K1, and GK (Santa Cruz Biotechnology, Santa Cruz, CA, USA); Vinculin (Sigma Chemicals); IRS-1, Shc 66 kDa, and p85 (Upstate Biotechnology Inc., Lake, Placid, NY, USA); and SREBP1 (Abcam, Paris, France). Anti-IR (B10) was a generous gift from Dr P. Gorden (National Institute of Diabetes and Digestive and Kidney Diseases, Bethesda, MD, USA). The signal was detected by enhanced chemiluminescence (ECL, GE Healthcare Europe GmbH, Saclay, France). The films were analyzed and signals quantified with the software MacBas V2.52 (Fuji PhotoFilm, USA, Inc.). Commercial antibodies directed against mammalian proteins used in western blot experiments have been previously shown to cross-react with chicken homolog protein at a good extent (confirming description of species specificity by trade companies, when available, see references from Dupont et al. and Tesseraud et al. quoted herein).

insulin-like growth factor1 (IGF1) blotting was performed as described previously (Beccavin et al. 1999) and IGF binding protein 1 band (28 kDa) was quantified using the PhosphoImager (Storm 840, GE Healthcare Europe GmbH, Saclay, France). PI3K activity was measured in p85 immuno-precipitates from liver and leg muscle homogenates (2 mg protein) using an anti-rat p85 subunit antibody (Upstate Biotechnology Inc, Euromedex, France) and protein G agarose beads (GE Healthcare Europe GmbH) as described previously (Dupont et al. 1998a).

RNA isolation and qRT-PCR analyses

After tissue powdering under liquid nitrogen, total RNA was extracted using RNase Mini or Midi kits (Qiagen). After RNase-free DNase treatment, RNA was reverse transcribed using random hexamers and SuperScriptII reverse transcriptase. Quantitative real-time PCRs (qRT-PCR) were performed in duplicate using an ABI Prism 7000 Sequence Detection System (Applied Biosystems, Foster City, CA, USA) (Pfaffl 2001). The sequences of primers (specifically designed or reproduced from the literature) are shown in Table 1. PCR product sequences were checked. For each gene assayed, experimental and reference samples were measured in one 96-well plate. Results were expressed as Δct (Δct = reference pool ct – sample ct); when negative, gene expression was lower in the sample than in the reference pool and the reverse when positive. Composition of the reference RNA pool is indicated in the footnotes to Table 3.

Statistical analysis

Data are presented as means ± s.e.m. Data were analyzed by ANOVA and treatment means were compared by Fisher’s test or t-test, significance was set at P<0.05.

Results

Metabolic parameters

Cumulative food intake (Table 2) was decreased at 1 h after a single anti-insulin serum injection (1hrfed-Ab versus 1hrfed-C) or at 5 h after three anti-insulin serum injections (5hrfed-Ab versus 5hrfed-C). Plasma glucose concentration was not significantly altered by food privation for 5 h (5hrfasted-C), whereas large hyperglycemia was induced by anti-insulin serum injection in the 1hrfed-Ab and 5hrfed-Ab groups when compared with their respective fed controls. Striking hyperglycemia was also present at 2 and 4 h before the second and third injections (not shown). Plasma NEFA levels significantly increased in the 5hrfed-Ab group, though at a lower extent than in 5hrfasted-C group. Plasma levels of αN2H2PN also significantly increased in the 5hrfed-Ab group, whereas they decreased following fasting (5hrfasted-C). Plasma glucagon was increased following fasting (5hrfasted-C group) and insulin privation (5hrfed-Ab); it is noteworthy that these increases have been underestimated (see footnote to Table 2). Plasma T3 was decreased after 5-h fasting (5hrfasted-C) and after 1- or 5-h insulin privation (1hrfed-Ab or 5hrfed-Ab).

IR signaling cascade

In the liver, anti-insulin serum significantly decreased tyrosine phosphorylation levels of IR β-subunit, IRS-1 and Shc 52 kDa isoform as well as PI3K activity at 1 h (1hrfed-Ab...
versus 1hrfed-C, Fig. 1A–D). The extent of these decreases is quite similar to that induced by 5-h fasting (5hrfasted-C versus 1hrfed-C). In the chicken tissues, Shc 52 kDa is the main isoform tyrosine phosphorylated in response to insulin (Dupont et al. 1998b). Phosphorylations of Akt, GSK3β, ERK2, and the S6 ribosomal protein were accordingly inhibited in both conditions (Fig. 1E–I). In chicken, only ERK2 form is phosphorylated (Duchêne et al. 2008a). In the muscle, in contrast to what was observed in the liver, injection of anti-insulin serum (1hrfed-Ab) or food deprivation for 5 h (5hrfasted-C) did not alter tyrosine phosphorylation levels of the IR-β subunit and IRS-1 (versus 1hrfed-C, Fig. 2A and B) or P38K activity (Fig. 2D). Both

Table 1 Primers used for quantitative RT-PCR of messengers in liver and/or leg muscles

<table>
<thead>
<tr>
<th>Messengers</th>
<th>Forward primer (5′–3′)</th>
<th>Reverse primer (5′–3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ribosomal 18S</td>
<td>Taqman control kit</td>
<td>Taqman control kit</td>
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<tr>
<td>β-Actin</td>
<td>CTGGCCACTAGCACAATGAAG</td>
<td>CTGGTTGCTGATCCACATCT</td>
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<tr>
<td>Transcription factors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Egr1</td>
<td>GACCCATCTGGCGCACACAA</td>
<td>AGCCCTGCTGCCATAGGTGG</td>
</tr>
<tr>
<td>PPARα</td>
<td>CAAACCAATACCTGAGATGG</td>
<td>CGAGGGTGATTCCTTGGAG</td>
</tr>
<tr>
<td>PPARγ</td>
<td>CAGGGAGGAAAGGCACGAAGA</td>
<td>TCCACAGGGCAGATCATC</td>
</tr>
<tr>
<td>SREBP1</td>
<td>GTCGCGGATCCGGGAAGA</td>
<td>CTCCCTGCAAGGCGCATC</td>
</tr>
<tr>
<td>THSRP2z (spot14)</td>
<td>TCTCCGGCAAGCGCAGAG</td>
<td>AAGACCCCTGCGACGAG</td>
</tr>
<tr>
<td>PGC1α</td>
<td>GGGACCGGTTGGAAGTGTG</td>
<td>GCCTGTGGACCTGGGAAG</td>
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</tbody>
</table>

Endocrine and signaling system

Table 2 Food intake and plasma metabolite and hormone parameters

<table>
<thead>
<tr>
<th>Group</th>
<th>1hrfed-C</th>
<th>5hrfed-C</th>
<th>1hrfed-Ab</th>
<th>5hrfed-Ab</th>
<th>5hrfasted-C</th>
<th>P level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Food intake, g</td>
<td>4.3 ± 0.3</td>
<td>18.4 ± 0.7</td>
<td>2.4 ± 0.4</td>
<td>14.0 ± 0.6</td>
<td>23.2 ± 0.7</td>
<td>0.0001</td>
</tr>
<tr>
<td>Glucose, mg/100 ml</td>
<td>264 ± 2</td>
<td>729 ± 6</td>
<td>434 ± 21</td>
<td>747 ± 28</td>
<td>232 ± 0.7</td>
<td>0.0001</td>
</tr>
<tr>
<td>NEFA, mg/100 ml</td>
<td>7.6 ± 0.3</td>
<td>7.3 ± 0.3</td>
<td>9.6 ± 0.6</td>
<td>11.6 ± 0.6</td>
<td>19.7 ± 1.7</td>
<td>0.0001</td>
</tr>
<tr>
<td>α-NH2NPN, mg/100 ml</td>
<td>93 ± 4</td>
<td>92 ± 6</td>
<td>106 ± 5</td>
<td>161 ± 10</td>
<td>65 ± 5</td>
<td>0.0001</td>
</tr>
<tr>
<td>Glucagon, ng/ml</td>
<td>0.12 ± 0.05</td>
<td>0.05 ± 0.02</td>
<td>0.39 ± 0.20</td>
<td>0.89 ± 0.20</td>
<td>0.89 ± 0.20</td>
<td>0.0001</td>
</tr>
<tr>
<td>T3, ng/ml</td>
<td>2.49 ± 0.23</td>
<td>2.19 ± 0.20</td>
<td>1.97 ± 0.16</td>
<td>1.06 ± 0.12</td>
<td>0.64 ± 0.08</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

*Food intake was 1-h cumulative intake for 1hrfed-C and 1hrfed-Ab groups following one normal serum injection or one anti-insulin serum injection respectively, and 5-h cumulative food intake for 1hrfed-C and 5hrfed-Ab groups following three injections of respective serum. Data are mean±s.e.m., n=7. For food intake, statistical analysis has been performed using two t-test comparing data at 1 or 5 h respectively; differences were significant at 1 h (P<0.008, a versus b) and 5 h (P<0.0001, A versus B). For plasma parameters, statistical analyses were performed including all experimental groups (Fα,30); values possessing different superscripts in the same row are significantly different (P<0.05). Free fatty acids (NEFA) are expressed as mg equivalent oleic acid/100 ml. α-NH2NPN (α-NH2 non-protein nitrogen) is expressed as mg equivalent leucine equivalent/100 ml. Glucagon mean values from the 1hrfed-Ab, 5hrfed-Ab, and 5hrfasted-C groups are indicated in parenthesis to mean that these values are underestimated; one value in the 1hrfed-Ab, two values in the 5hrfed-Ab group, and two values in the 5hrfasted-C group were out of range and replaced by 1.5 ng/ml, i.e. the maximum reliable value in the assay. Paucity of plasma did not permit further measurements at lower plasma concentrations.

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Figure 1  Insulin signaling in chicken liver 1 h after one anti-insulin serum injection. It shows western blots (A) phospho-IR β, (B) IRS-1, and (C) Shc 52 kDa on tyrosine residues and (E) phospho-Akt, (F) GSK3 β, (G) ERK2, (H) p70S6 kinase, and (I) S6 ribosomal protein) and (D) PI3K activities in liver lysates of fed chickens injected with either normal serum (1hrfed-C) or anti-insulin serum (1hrfed-Ab). As an additional control, one group of chicken fasted for 5 h received three injections of normal serum (5hrfasted-C). In western blots, when immuno-precipitations (IP) had been performed before gel electrophoresis, the antibody used is indicated as: IP:xxx; immune sera used to reveal protein phosphorylation levels are indicated to the left of gels (for instance, PY20 is directed against anti-phosphotyrosine residues). The levels of phosphorylation of IR, IRS1, Akt, and ERK2 have been normalized by the level of the corresponding protein using specific antibodies indicated to the left of gels. For Shc, the phosphorylation of 52 kDa isoform (the main tyrosine phosphorylated Shc isoform in response to insulin) has been normalized by the level of 66 kDa isoform. Phosphorylation levels of GSK3 and S6 ribosomal protein have been normalized by the level of vinculin. In chicken tissues, only ERK2 is phosphorylated. For each protein, gels show bands, which have been underlined per group with three chickens per group. From the left to the right, groups are 1hrfed-C, 1hrfed-Ab, and 5hrfasted-C groups. Below each gel, histograms represent mean ± S.E.M from two gels, with n = 6/group (the legend for groups is on top of the figure). When letters above bars differ, means significantly differ by at least P<0.05.
Figure 2  Insulin signaling in chicken leg muscles 1 h after one anti-insulin serum injection. It shows western blots ((A) phospho-IR β, (B) IRS-1, and (C) Shc 52 kDa on tyrosine residues and (E) phospho-Akt, (F) GSK3 β, (G) ERK2, (H) p70S6 kinase, and (I) S6 ribosomal protein) and (D) PI3′-kinase activities in leg muscle lysates of fed chickens injected with either normal serum (1hrfed-C) or anti-insulin serum (1hrfed-Ab). As an additional control, one group of chicken fasted for 5 h received three injections of normal serum (5hrfasted-C). In western blots, antibodies used for immuno-precipitations (IP), measurements of phosphorylations and normalizations were the same as indicated in the legend to Fig. 1. Phosphorylation of T389 and T421/424 residues of P70-S6 kinase has been measured using specific antibodies indicated to the left of gels. For each protein, gels show bands, which have been underlined per group with three chickens per group. From the left to the right, groups are 1hrfed-C, 1hrfed-Ab, and 5hrfasted-C groups. Below each gel, histograms represent mean ± S.E.M., from two gels with n=6/group (the legend for groups is on top of the figure). When letters above bars differ, means significantly differ by at least P<0.05.
conditions, however, decreased tyrosine phosphorylation of the Shc 52 kDa protein (Fig. 2C). In addition, phosphorylations of Akt, GSK3β, ERK2, P70S6K1, and S6 were all decreased by insulin privation or 5-h fasting (Fig. 2E–I). In both liver and muscle, AMPK phosphorylation on T172 residue in the α isozyme was unaltered by any of the present experimental conditions, meaning that AMPK was not activated (data not shown).

Effects of insulin privation on gene expression and protein levels

We next investigated whether insulin privation could alter expression of some specific genes in the liver and the muscle of fed chickens. Genes were chosen as insulin sensitive in mammals and, for some, as potentially invariant. Selected genes are involved in transcription, endocrine systems, and metabolism. Quantitative RT-PCR was performed at 5 h, comparing three conditions (5hrfed-Ab, 5hrfed-C, and 5hrfasted-C). In the liver (Table 3), 18S and β-actin mRNA were found invariant in the three experimental conditions. Expression of Egr1 was significantly reduced by anti-insulin serum but not by fasting. Expression of three transcription factors (PPARγ, SREBP1, and THRSPα) was significantly decreased by anti-insulin serum or fasting. Both anti-insulin serum and fasting also decreased the expression of D2-deiodinase, glucokinase (GK), and the fatty acid synthase (FASN) genes, whereas these two conditions increased the abundance of D3-deiodinase and IGFBP1 mRNA. In contrast, several other mRNA (PPARα, PPARβ, adiponectin, adiponectin receptor R1, GSK3β, Cox4, and cathepsin B) were not significantly modified by either treatment (data not shown). In the muscle, only 6 out of the 20 genes studied were altered by anti-insulin antibodies, fasting, or both conditions (Table 4). The mRNA for PPARβ and atrogin-1 were increased by anti-insulin serum or fasting. Gene expression for avUCP and MURF1 were increased, only by fasting. The expression of D3-deiodinase gene was also significantly decreased by fasting. The Egr1 mRNA level was selectively decreased by anti-insulin antibodies. Expression of several other genes (18S, β-actin, PGC1α, PPARα, PPARγ, SREBP1, THRSPα, D2-deiodinase, adiponectin, adiponectin receptor R1, GSK3β, Cox4, and cathepsin B) were not altered.

Finally, the levels of some proteins have been assessed. Liver GK and SREBP1 protein decreased at 5 h following insulin privation or fasting (Fig. 3A and B). The plasma IGFBP1 protein increased following insulin privation or fasting (5hfed-Ab or 5hrfasted-C, Fig. 3C).

Discussion

Privation of insulin by immuno-neutralization rapidly induced large hyperglycemia in fed chickens. Lipid and amino acid metabolism was also disturbed. The increase in plasma NEFA most likely resulted from the development of an early hyper-glucagonemia (glucagon is the lipolytic hormone in birds while no anti-lipolytic effect has been demonstrated for insulin in chicken adipocytes (see references in (Simón 1989))). The fact that glucose requires the presence of insulin to inhibit pancreatic glucagon release has been previously recognized in ducks first, and then in mammals (Miahle’s group references in Simón 1989). The increase in total plasma amino acids following insulin privation may have

### Table 3 Liver genes altered at 5 h by insulin antibody or fasting (Δct)*

<table>
<thead>
<tr>
<th>Gene name</th>
<th>5hrfed-C</th>
<th>5hrfed-Ab</th>
<th>5hrfasted-C</th>
<th>P level, ANOVA</th>
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<tbody>
<tr>
<td><strong>Transcription factors</strong></td>
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<td></td>
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</tr>
<tr>
<td>Egr1</td>
<td>-2.4 ± 0.2</td>
<td>-4.2 ± 0.3</td>
<td>-2.9 ± 0.2</td>
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<td>PPARγ</td>
<td>0.15 ± 0.2</td>
<td>(-0.8 ± 0.2)</td>
<td>-0.7 ± 0.3</td>
<td>0.03</td>
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<td>SREBP1</td>
<td>-0.4 ± 0.3</td>
<td>-1.7 ± 0.5</td>
<td>-1.8 ± 0.3</td>
<td>0.05</td>
</tr>
<tr>
<td>THRSPα</td>
<td>1.2 ± 0.2</td>
<td>-0.9 ± 0.6</td>
<td>-1.4 ± 0.2</td>
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<td><strong>Endocrine system</strong></td>
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<td>D2-Deiodinase</td>
<td>1.3 ± 0.3</td>
<td>-2.1 ± 1.1</td>
<td>-3.4 ± 0.6</td>
<td>0.001</td>
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<tr>
<td>D3-Deiodinase</td>
<td>-1.5 ± 0.6</td>
<td>0.8 ± 0.8</td>
<td>3.1 ± 0.7</td>
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<td>IGFBP1</td>
<td>-1.3 ± 0.2</td>
<td>1.7 ± 0.5</td>
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</tr>
<tr>
<td>GK</td>
<td>0.72 ± 0.29</td>
<td>-0.02 ± 0.21</td>
<td>-0.24 ± 0.06</td>
<td>0.01</td>
</tr>
<tr>
<td>FASN</td>
<td>0.2 ± 0.3</td>
<td>-1.4 ± 0.8</td>
<td>-2.3 ± 0.3</td>
<td>0.01</td>
</tr>
</tbody>
</table>

* mRNA levels were determined by qRT-PCR analysis using primers listed in Table 1. A RNA pool was prepared from 15 aliquots of 10 micrograms issued from one chicken each of the five experimental groups and three tissues: liver, leg, and breast muscles and used as a reference standard. Data were calculated as Δct (Δct = RNA reference pool ct—sample ct) for each messenger assayed at 1/100 dilution, except for: 18S and β-actin (assayed at 1/5000 dilution) and GK (assayed at 1/8 dilution). When Δct is negative, messenger is less expressed in the sample than in the pool reference and the reverse is true when Δct is positive (so, the higher Δct value is, the higher the mRNA abundance is). Data are presented as mean ± S.D. for n=7. ANOVA significance P levels are indicated in the last column to the right. One bird from the 5hrfed-Ab group exhibited expression level largely out of range of this group for several genes. For PPARγ transcript, the value of this chicken (Δct = 4.0 ± 0.69) was excluded from the analyses. For this reason, the mean for n=6 is indicated in parenthesis. For each gene mRNA, means possessing different superscripts are significantly different. Several mRNAs were not significantly altered in the liver by the experimental treatments: ribosomal 18S, β-Actin, PPARα, PPARβ, adiponectin, adiponectin R2, D1-deiodinase, and Cox4.

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Table 4  Leg muscle genes altered at 5 h by insulin antibody or fasting (Δct)*

<table>
<thead>
<tr>
<th>Gene name</th>
<th>5hrfed-C</th>
<th>5hrfed-Ab</th>
<th>5hrfasted-C</th>
<th>P level, ANOVA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eg1</td>
<td>1.3±0.8</td>
<td>0.3b±0.5</td>
<td>1.1±0.8</td>
<td>0.03</td>
</tr>
<tr>
<td>PPARβ6</td>
<td>0.5±0.1</td>
<td>0.3±0.2</td>
<td>0.5±0.1</td>
<td>0.002</td>
</tr>
<tr>
<td>D3-Deiodinase</td>
<td>1.3±0.4</td>
<td>-1.5±0.2</td>
<td>-2.4b±0.2</td>
<td>0.04</td>
</tr>
<tr>
<td>Metabolism</td>
<td>0.6±0.3</td>
<td>1.3±0.2</td>
<td>2.4±0.3</td>
<td>0.0003</td>
</tr>
<tr>
<td>AvUCP</td>
<td>-1.2±0.4</td>
<td>-0.3b±0.3</td>
<td>1.8±0.3</td>
<td>0.0001</td>
</tr>
<tr>
<td>Atrogin-1</td>
<td>-0.4±0.3</td>
<td>0.1h±0.2</td>
<td>1.6±0.3</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

*Expression levels were determined by qRT-PCR analysis using primers listed in Table 1 and the RNA reference pool described in Table 3. Data were calculated as Δct (Δct = pool reference ct – sample ct) for each mRNA assayed at 1/100 dilution, except for 18S and β-actin messengers which were assayed at 1/5000 dilution. When Δct is negative, mRNA levels are lower in the sample than in the pool reference and the reverse is true when Δct was positive (so, the higher Δct value is, the higher the mRNA abundance is). Data are presented as mean ± S.E.M. for n = 7. ANOVA P levels of significance are indicated in the last column to the right. For each gene mRNA, means possessing different superscripts are significantly different. Several transcripts were not significantly altered by treatments: 18S, Metabolism genes have been assessed in the liver and the muscle.

In the chicken liver, the main steps of the insulin signaling cascade are, accordingly, dependent on the insulin status (Simon 1989, Tesseraud et al. 2007b). A rapid decrease in food intake was the last general perturbation observed, which suggests that glucose and insulin interact in a complex manner to balance orexigenic and anorexigenic pathways in the brain and/or at the peripheral tissues. Such impairment in food intake had not been previously observed most likely because hyperglycemia was not so high in comparison with the present results (Simon et al. 2000). One possible hypothesis could be that in the present conditions, cerebrospinal fluid concentration of insulin had not yet changed whereas that of glucose increased, leading to a satiety signal. In order to get some insight into the mechanisms involved into the development of such acute `diabetic' status, the insulin signaling pathways and the expression of some specific genes have been assessed in the liver and the muscle.

In the chicken liver, the main steps of the insulin signaling cascade are, accordingly, dependent on the insulin status (Dupont et al. 1998b,c, 2004). In the present study, a fasting period as short as 5 h appeared sufficient to turn-off liver insulin signals. IR, IRS-1, and Shc tyrosine phosphorylation; PI3K activity; and phosphorylations of Akt, GSK3, ERK2, and S6 ribosomal protein were all strongly inhibited, clearly indicating the development of a catabolic status. These changes were large and very fast, which may result from high internal body temperature (42°C) and turnover rate of chicken metabolism. Insulin privation in fed chicken also turned off all the same signaling components in the liver at 1 h to an extent similar to that induced by 5 h of fasting.

In contrast, in leg muscles, IR and IRS-1 tyrosine phosphorylation and PI3K activity were totally unresponsive to fasting or insulin privation. However, Shc 52 kDa tyrosine phosphorylation, thr/ser phosphorylation of P70S6K1, and phosphorylation of the other proteins assessed in the liver, all decreased in both situations (5-h fasting and 1-h insulin privation). As previously observed (Dupont et al. 2004), high amounts of p85 subunit of PI3K and high activities of PI3K are constitutively present in chicken muscle when compared with rat muscle. In normal or diabetic mice, a decrease in the amount of PI3K does improve insulin sensitivity (Terauchi et al. 1999, Ueki et al. 2002). Recently, it has been shown that an excess of p85 may impair insulin signaling by competing with binding of the active p85–p110 complex to IRS1 and activating two counter-regulatory enzymes: PTEN (lipid phosphatase and Tensin homolog) and c-Jun N-terminal kinase (JNK; Taniguchi et al. 2006a,b, 2007). The presence of high amounts of p85 subunit in chicken muscle may therefore contribute to the low sensitivity to exogenous insulin observed in chicken. Other surprising features for insulin signal cascade in chicken muscle were largely substantiated and extended in the present study. Namely, several components generally considered as downstream effectors of PI3K (Akt, GSK3β, and P70S6K1) up to the subsequent phosphorylation of S6 ribosomal protein were inactivated by insulin privation in fed chickens (and 5 h fasting). Amino acids have been shown to activate P70S6K1 in vitro, mimicking insulin action (Tesseraud et al. 2007a). In the present conditions, following insulin privation, amino acid levels increased in plasma but were ineffective at supporting an active P70S6K1. The alteration observed for the phosphorylation of ERK2 could be related to the decrease in Shc tyrosine phosphorylation. IRS-2, another important IR substrate in mammals, has not yet been characterized in chicken to our knowledge. An IRS-2 homolog coding sequence has been recently suggested in the chicken genome on chromosome 1 (Ensemble release 47, October 2007 at http://www.ensembl.org/Gallus_gallus/index.html). However, it is hard to conceive that Shc and/or another IRS would account for insulin action in chicken muscle, via the typical PI3K pathway described in mammals (White & Kahn 1994). Glucagon receptors activate hydrolysis of phosphatidylinositol phosphates in addition to the activation of the major adenylyl
cyclase/protein kinase A signaling pathway (Unson et al. 1989). No evidence of an alternative pathway has been clearly supported yet for insulin receptors. A unifying hypothesis would be that transient or discrete changes in IR tyrosine phosphorylation, undetectable with the present methodology may occur in vivo, which would account for small changes in Shc 52 kDa tyrosine phosphorylation as presently and previously (Dupont et al. 2004) observed in chicken muscle. Within this hypothesis, Shc and potentially other IRS, but not IRS-1, would control signaling steps beyond the PI3K through an unknown and hypothetical bypass or parallel mechanism. This would enable insulin to exert some control in chicken muscle.

It is also noteworthy that AMPK has not been activated in the liver or the muscle by either 1-h insulin privation or 5-h fasting. AMPK, a serine–threonine kinase now considered as an energy sensor, is activated in response to nutrient deficiency or stress to support cellular metabolism and prevent a decrease in ATP cellular level (Long & Zierat 2006). Present results suggest that AMPK is not immediately recruited in case of metabolic perturbations (including in muscle and despite the large increase in plasma NEFA levels). This conclusion is also supported by the absence of AMPK activation in chicken brain, liver, or skeletal muscle in response to a 24-h fast; AMPK was, however, activated in muscle after a 48-h fast (Prosakowiec-Wegalz et al. 2006). In contrast, in the ovary of several species, including hen, AMPK activity varies during follicle development, independently of nutrition level (Dupont et al. 2008). In the present study, it was of interest to investigate whether the expression of some genes and proteins, known as insulin sensitive in mammals, had been altered in the liver or the muscle following insulin privation (5hrfed-Ab).

In the liver, mRNA levels of several transcription factors were altered by insulin privation. Egr1 (a zinc finger protein) is an immediate early transcription factor expressed in multiple tissues and cell types (Sukhatme et al. 1987). It was selectively decreased in the ‘diabetic’ status at 5 h but not by...
the 5-h fasting period. Interestingly, turning off the insulin cascade is not sufficient to decrease Egr1 expression since fasting did turn the cascade off without inhibiting Egr1 expression in the present conditions, which suggests a complex control of Egr1 expression. This, however, further supports the insulin specificity for the control of Egr1, which is a vital transcription factor for many genes. Several studies implicate Egr1 in the control of hepatic malic enzyme and A1 apolipoprotein genes in rat and human cells respectively (Kilbourne et al. 1990, Barroso et al. 1999). However, in the present study, changes in Egr1 messenger are most likely not involved in the changes observed for the lipogenic pathway (see next paragraph), since the lipogenic pathway was altered after both insulin privation or fasting.

Insulin privation and 5-h fasting decreased three other transcription factors in the liver, namely PPARγ, SREBP1, and THRSPζ (Spot14), to about the same extent. Other PPAR family members were unchanged. Most likely, the changes in SREBP1 and PPARγ were coordinated, and accounted for the inhibition of FASN gene expression. Liver is the lipogenic organ in chickens (Simon 1989) and SREBP1 control of lipogenesis has been clearly demonstrated (Gondret et al. 2001). Importantly, both insulin privation and fasting were able to rapidly decrease the total amount of the SREBP1 protein, most likely accounting for the decrease in PPARγ expression. The present results on SREBP1 messenger and protein following insulin privation are the mirror image of changes observed in 24-h fasted–5-h re-fed chickens (Zhang & Hillgartner 2004). Whether THRSPζ expression is also dependent on SREBP1 is not yet clear. In addition to control by other transcriptional factors, THRSPζ mRNA levels are regulated by T3 and, most likely, by pancreatic hormones in the present study and in the rat (Kinlaw et al. 1987). In the liver, THRSPζ is differentially expressed in chicken lines exhibiting large differences in fat content and is dependent on the thyroid hormone status (Wang et al. 2007a,b). It is also present in adipose tissue (Wang et al. 2007a,b), though chicken adipocytes are not considered as active lipogenic sites (Simon 1989). The decrease in liver THRSPζ messenger is very likely also to contribute to the inhibition of FASN gene expression since this factor is involved in the control of lipogenic enzymes (Roder et al. 2000). Rat hepatocytes, transfected with a spot 14 antisense oligonucleotide, express decreased mRNA levels in enzymes involved in the lipogenic pathway, including FASN (Kinlaw et al. 1995).

Insulin privation (and fasting) disturbed several components of the endocrine or metabolic systems. Deiodinases are seleno–enzymes that regulate T3 availability in peripheral tissues. Among them, liver D2-deiodinase expression was decreased while D3-deiodinase expression was increased, which would account for the reduction in plasma T3. Such concomitant perturbations should rapidly inhibit the lipogenic pathway, since a potent and positive synergy occurs between T3 and insulin at several steps (references in Simon 1989). The present results suggest that the fine tuning of thyroid hormone metabolism is also insulin and most likely glucagon dependent in addition to other endocrine and nutritional controls (Darras et al. 2006). The hepatic adiponectin system (R2 and ligand) remained unaltered by insulin privation. In contrast, liver IGFBP1 mRNA clearly increased following insulin privation or 5-h fast. Therefore, rapid liver IGFBP1 synthesis should contribute to the increase in plasma IGFBP1 protein. Present and previous (Beccavin et al. 1999) observations make this protein a typical IGFBP1 protein, which is increased in mammals as a complication of diabetes. Under many circumstances, IGFBP-1 is inversely related to insulin concentration in human. In most strains of IGFBP-1 transgenic mice, fasting hyperglycemia, impaired glucose tolerance, and a modest insulin resistance in skeletal muscle and liver develop (Rajkumar et al. 1996).

In mammals, one of the major roles of insulin in the liver is stimulation of the synthesis and activity of GK (Narkewicz et al. 1990). The existence of a typical GK and the nutritional regulation of its activity have only recently been demonstrated in chicken liver (Berradi et al. 2005). The present results show the insulin dependency of GK mRNA in the fed chicken liver and suggest a rapid glucagon inhibitory effect in the fasting state. In the absence of a functional cytosolic PEPCK activity in chicken liver (Tinker et al. 1983, i.e. the regulatory component of gluconeogenesis in mammals), the decrease in liver GK mRNA and protein in fed chicken following insulin privation should decrease liver glucose utilization and contribute to the development of marked hyperglycemia. Indeed, it has been shown that GK knockout mice display mild basal hyperglycemia and exhibit a profound defect in glucose turnover (Postic et al. 1999).

In leg muscles, insulin privation altered only a few of the gene considered in the present study. As in the liver, muscle Egr1 mRNA was decreased by insulin privation at 5 h and not by fasting. PPARβ and atrogin-1 mRNAs were increased by insulin privation but also by fasting. In another study, atrogin-1 mRNA increased then decreased in chicken muscle along with a 24-h fasting–2-h re-feeding experiment, most likely under the control of the Akt and FOXO-1 pathways (Nakashima et al. 2006). In a quail fibroblast cell line (QT6 cells), atrogin-1 transcript was inhibited at 5 h following insulin exposure with an early stimulation of Akt phosphorylation (Tesseraud et al. 2007c). Herein, fasting modified the expression of MURF1 and avUCP in muscle. Both atrogin-1 and MURF1 are muscle-specific atrogenes involved in the regulation of ubiquitin/proteasome-mediated proteolysis (Lecker 2003). This suggests that fasting rapidly induces proteolysis in chicken muscle. In the fed state, insulin would stimulate muscle protein accretion by inhibiting proteolysis through inhibition of atrogin-1 expression, in addition to stimulating cell amino transport and protein synthesis. Expression of other transcription factors (PGC1α, PPARβ, PPARγ, SREBP1, and THRSPζ) was not altered in muscle following insulin privation. In the present study, insulin privation induced a large increase in plasma NEFA, albeit the abundance of avUCP mRNA in muscle was not affected. Therefore, the large increase in muscle avUCP mRNA...
induced by the 5-h fast might rely only partly on the increase in fatty acid availability. Until now, avUCP (the unique UCP thus far identified in chicken) has been involved in thermogenesis, fatty acid transport, and control of superoxide production in muscle mitochondria (Collin et al. 2005, Abe et al. 2006).

In mammals, skeletal muscles clear large amounts of plasma glucose during anabolic phases (DeFronzo et al. 1992). The existence of typical Glut4 glucose transporters is still under debate in chickens and recently in sparrows (Sweeza & Braun 2006), which could come from the lack of cross-reactivity of antibodies directed against mammalian Glut4s. The existence of a functional and Glut4-related isoform of transporter has been suggested only in ducks (Thomas-Delloye et al. 1999). A Glut4 homolog has not been identified in the chicken genome; however, the chicken genome has not been completely sequenced yet. To date, only Glut1, Glut2, Glut3, and Glut8 homologs from the 13 Glut isoforms found in mammals (Joost et al. 2002) have been identified in chickens (Seki et al. 2003). The existence of a functional insulin-dependent glucose transporter in chicken muscle has been strongly suggested in vivo using 2-deoxyglucose uptake following insulin injection, at least in some muscles (Nishiki et al. 2008, Tokushima et al. 2003). The large hyperglycemia observed after insulin privation in fed chickens in the present study also favors the existence of functional Glut4-related glucose transporters in chicken.

In conclusion, insulin signaling is peculiar in chicken muscle. An acute ‘diabetic’ status develops rapidly after insulin immunono-neutralization in fed chickens. Hyper-glucagonemia and low T₄ are early hormonal perturbations in the absence of insulin signal. Expression of several genes involved in regulatory pathways or metabolisms appeared clearly insulin dependent in the liver and the muscle, evidencing pleiotropic effects of insulin in fed chicken.

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