DIRECT RECRUITMENT OF INSULIN RECEPTOR AND ERK SIGNALING CASCADE TO INSULIN-INDUCIBLE GENE LOCI

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OBJECTIVE—Insulin receptor (IR) translocates to the nucleus, but its recruitment to gene loci has not been demonstrated. Here, we tested the hypothesis that IR and its downstream mitogenic transducers are corecruited to two prototypic insulin-inducible genes: early growth response 1 (egr-1), involved in mitogenic response, and glucokinase (Gck), encoding a key metabolic enzyme.

RESEARCH DESIGN AND METHODS—We used RNA and chromatin from insulin-treated rat hepatic tumor cell line expressing human insulin receptor (HTC-IR) and livers from lean and insulin-resistant ob/ob glucose-fed mice in quantitative RT-PCR and chromatin immunoprecipitation studies to determine gene expression levels and associated recruitment of RNA polymerase II (Pol II), insulin receptor, and cognate signaling proteins to gene loci, respectively.

RESULTS—Insulin-induced egr-1 mRNA in HTC-IR cells was associated with corecruitment of IR signaling cascade (IR, SOS, Grb2, B-Raf, MEK, and ERK) to this gene. Recruitment profiles of phosphorylated IR, B-Raf, MEK, and Erk along egr-1 transcribed region were similar to those of elongating Pol II. Glucose-feeding increased Gck mRNA expression in livers of lean but not ob/ob mice. In lean mice, there was glucose feeding-induced recruitment of IR and its transducers to Gck gene synchronized with elongating Pol II. In sharp contrast, in glucose-fed ob/ob mice, the Gck recruitment patterns of active MEK/Erk, IR, and Pol II were asynchronous.

CONCLUSIONS—IR and its signal transducers recruited to genes coupled to elongating Pol II may play a role in maintaining productive mRNA synthesis of target genes. These studies suggest a possibility that impaired Pol II processivity along genes bearing aberrant levels of IR/signal transducers is a previously unrecognized facet of insulin resistance. Diabetes 60:127–137, 2011

The insulin receptor (IR) is a member of a family of receptor tyrosine kinases (RTKs) that include the epidermal growth factor receptor, fibroblast growth factor receptor, and several others (1). Studies have shown that RTKs are translocated to the nucleus upon stimulation, including fibroblast growth factor receptor (2,3) and epidermal growth factor receptor, and several others (1).

Studies have shown that RTKs are translocated to the nucleus upon stimulation, including fibroblast growth factor receptor (2,3) and epidermal growth factor receptor (4). Importantly, there is evidence for recruitment of RTKs to chromatin and gene loci, and for several RTKs the recruitment increases transcription (2,5,6,7). RTKs are not the only kinases found along genes. For instance, active ERK1/2, MEK, p38, and AMPK are also recruited to genes (8,9,10,11).

IR also translocates to the nucleus after binding its ligand (12,13,14). For instance, mice fed a glucose meal showed an increase in nuclear IR, which correlated with the glucose-stimulated rise of serum insulin (13,15). The fact that RTKs and the terminal mitogen-activated protein kinases (MAPKs) can be inducibly recruited to chromatin presents the possibility that canonical insulin signaling pathways are recapitulated along gene loci. Here, we provide the first evidence that not only the IR but also most of the ERK cascade components, as well as ERK dual-specificity phosphatase (16), are corecruited to inducible genes.

RESEARCH DESIGN AND METHODS

Reagents. BSA (cat. no. A2153), salmon sperm DNA (cat. no. D1626), and protein A (cat. no. F7837) were from Sigma, protease K was from Invitrogen (cat. no. 25530-015), and Humulin N was from Eli Lilly. Matrix chromatin immunoprecipitation (ChIP) 96-well polystyrene plates were from Corning (Costar cat. no. 9018), and polypropylene plates were from Bioexpress (T-3060).

Tissue culture and insulin treatment. HTC and HTC-IR cells were maintained and insulin treated as in ref (17) with the exception that 1 × 10⁻⁸ M insulin was used.

Mice and glucose feeding experiments. Male C57BL/6 and ob/ob mice were purchased from The Jackson Laboratory (Bar Harbor, ME) and were maintained, glucose fed, and killed as described previously (13). Blood was collected just prior to sacrifice. Livers were removed after whole-animal perfusion with sterile, cold PBS and flash frozen in liquid nitrogen. Blood glucose was measured using One Touch Ultra system (LifeScan Inc.). Plasma insulin levels were measured using the Linco insulin ELISA (Millipore). All procedures were done in accordance with current National Institutes of Health guidelines and approved by the Animal Care and Use Committee of the University of Washington.

RNA extraction and cDNA synthesis. RNA was extracted from cell pellets or tissue fragments using Trizol reagent as per the manufacturer’s protocol. To synthesize cDNA, 400 ng of Trizol extracted total RNA was used in reverse transcription reactions with 200 units of MMLV reverse transcriptase (Invitrogen) and random hexamers. RT reactions were diluted 100-fold prior to running quantitative PCR (18).

Matrix ChIP. Chromatin shearing was done using either Misonix 3000 microprobe (1 ml IP buffer [8], six rounds of sonication power 5, 15 s, on ice) or Diagenode Bioruptor (100 μl IP buffer, 30 rounds 30 s ON/30 s OFF, high power, 4°C). The suspension was cleared by centrifugation at 12,000g (10 min at 4°C), and the supernatant, representing sheared chromatin, was aliquoted and stored at −80°C. ChIP assays were done using the Matrix ChIP platform in 96-well microplates as described before (8,19). ChIP DNA samples were assayed by quantitative PCR. PCR calibration curves were generated for each primer pair from a dilution series of total rat or mouse sheared genomic DNA. The PCR primer efficiency curve was fit to cycle threshold versus log (genomic DNA dilutions) using an r² best fit. DNA concentration values for each ChIP and input DNA samples were calculated from their respective average cycle threshold values. Final results are expressed as a fraction of input DNA (8). Matrix ChIP PCR primers are shown in supplementary Table 1 and the list of antibodies are shown in supplementary Table 2, available in an online appendix at http://diabetes.diabetesjournals.org/cgi/content/full/db09-1806/DC1.
RESULTS

Insulin stimulates egr-1 transcription in hepatocyte culture. We examined expression of the mitogen-sensitive immediate early gene, early growth response 1 (egr-1), as a model of insulin-induced transcription in rat HTC-IR cells overexpressing human IR (20). Figure 1A (left) shows that the egr-1 transcript increased by >10-fold after 60 min of insulin treatment. β-actin transcript levels did not change (Fig. 1A, right).

To determine whether the increase in egr-1 transcript was due to increased transcription, Matrix ChIP (8) for polymerase II (Pol II) was used with primers to or near the egr-1 locus. Pol II density increased within the body of the gene beginning as early as 5 min after treatment (Fig. 1C). Peak Pol II binding at the end of egr-1 coincided with the beginning of the increase in its transcript, 10 min after insulin treatment (Fig. 1A, left panel), suggesting that transcription upregulation was at least partly responsible for the induction of egr-1 transcript.

Insulin induces recruitment of the IR transducers to the egr-1 locus in hepatocyte culture. We used antibodies to the β-subunit of IR with Matrix ChIP to determine whether IR was recruited to egr-1 in insulin-treated HTC-IR cells. IR was recruited to egr-1 within the transcribed region, beginning 5 min after insulin treatment and remaining at the same level for 15 min (Fig. 2C, row 1).

To determine whether the recruited IR was activated, we used Matrix ChIP with an antibody to IR phosphorylated at tyrosine 1146, a site autophosphorylated upon binding of the receptor to insulin (21). Interestingly, phospho-IR had a large transient increase at both sites within the coding region of egr-1, with kinetics similar to those of Pol II (Fig. 2C, row 2). The insulin-induced total and phospho-IR ChIP signals were much less intense in the rat parental HTC cell line without human IR (20), while the histone H3 lysine 27 trimethylation, H3K27m3, ChIP signals were the same (supplementary Fig. 1, available in an online appendix). These results confirm that the antibodies detect total and phospho-IR. The more robust insulin-induced activation of IR at egr-1 in HTC-IR cells paralleled higher levels of recruitment of Pol II and phospho-Erk (supplementary Fig. 1), suggesting that these insulin-induced changes were linked.

Upon activation and autophosphorylation of IR, the insulin receptor substrates (IRSs) and Src homology 2 domain containing protein (Shc) are recruited to phosphotyrosines on the receptor and then become phosphorylated at tyrosines themselves (21,22). These IRSs recruit several proteins including the transducing adapter, Grb2 (22). Grb2 in turn recruits the guanine-nucleotide exchange factor, SOS (21,23), which then activates Ras (22,24). Several of these transducers including IRS-1 (25),...
Shc (26), and Grb2 (27) have been found in the nucleus. Thus, antibodies to IRS-1, Shc, Grb2, SOS, and Ras were used next in Matrix ChIP to determine which if any of these components of the insulin receptor complex were recruited to \( egr-1 \). We observed recruitment of Grb2 and SOS but not IRS-1, Shc, or Ras to the \( egr-1 \) gene (Fig. 2C, rows 3 and 4). These data are the first example of corecruitment of the IR and downstream signaling components to a gene locus and, importantly, in a manner associated with activation of the gene by insulin. The failure to detect IRS-1, Shc, and Ras may reflect either their absence at the \( egr-1 \) locus or that the antibodies used do not work in ChIP assays.

**Insulin induces the recruitment and activation of the Raf–MEK–ERK MAPK pathway at the \( egr-1 \) locus in hepatocyte culture.** The fact that we were able to demonstrate recruitment of SOS to \( egr-1 \) and that SOS activates the ERK MAPK pathway through the small GTP-binding protein, Ras, led us to probe recruitment of the Raf–MEK–ERK pathway to \( egr-1 \). We used antibodies to B-Raf phosphorylated at T598 and S601 (pB-Raf) (28), MEK1/2 phosphorylated at S217/221 (pMEK), and ERK1/2 phosphorylated at S38/56 (pERK).
phosphorylated at T202 and Y204 (pERK) in Matrix ChIP. All three phosphorylated kinases were transiently but robustly recruited to both sites within the transcribed region of *egr-1* and with kinetics that were very similar to those of the phosphorylated IR (Fig. 3B, rows 2, 4, and 6). ChIP with antibodies to the total kinases, however, showed less pronounced and less transient patterns (Fig. 3B, rows 1, 3, and 5).

The recruitment of insulin-induced signaling proteins to an insulin-responsive gene suggests their involvement in transcription. To examine the relationship between transcribing Pol II and locus-bound kinases, we compared the recruitment kinetics of the total and activated signaling proteins with those of Pol II within the coding regions of *egr-1*. To normalize the differing efficiencies of pulldown for each antibody, we plotted the ratio of each protein’s signal at each time point versus the signal at the last time point (supplementary Fig. 2, available in an online appendix). The nearly identical binding profile of active IRβ, kinases, and Pol II suggests that, after the recruitment of transcription.
IR and MAPK signaling components, the activation/deactivation of the kinases is tightly coupled to IR, a chain of events that are linked to RNA Pol II elongation. Compared with the insulin-induced levels of binding of total kinases, the phospho-kinase signals exhibited greater inducibility and were short-lived (supplementary Fig. 2). The transient nature of the phospho-kinase signals may reflect rapid dephosphorylation by phosphatases that are recruited to egr-1 in response to insulin. MKP-1 is a member of the dual-specificity family of phosphatases that inactivate Erk1/2 through dephosphorylation of its pThr202 and pTyr204 residues (29) (Fig. 3A). MKP-1 is insulin-responsive (30,31) and is found in the nucleus (32,33). ChiP assays for MKP-1 revealed insulin-induced recruitment to egr-1 (Fig. 3B, row 7). This, together with the corresponding decrease in phospho-ERK, but not total ERK (5–10 min, 232-bp site), suggests the possibility that Erk1/2 found at the egr-1 locus is inactivated in situ and that at least a fraction of the dephosphorylated Erk1/2 remains bound to the gene.

The serum responsive factor activates egr-1 by binding to several promoter elements (34). Matrix ChiP with an antibody to serum responsive factor showed a high constitutive and a small insulin-inducible component at the −252 bp and +232 bp sites and no binding at the end of the gene (Fig. 3, row 8). Thus, the trend toward higher levels at the transcribed region of egr-1 was specific to the IR signaling and MAPK proteins (Fig. 3B). Together, the above evidence is the first to suggest the recruitment of an entire activated MAPK pathway as well as cognate phosphatase at a gene in mammalian cells across the transcribed region.

**Increased recruitment of IR and the MAPK signaling module to the Gck gene in livers of glucose-fed mice.** Glucose feeding of fasted mice was previously shown to cause a rapid increase in serum glucose, serum insulin, and nuclear translocation of IR (13). In the following experiments, fasted C57BL/6 mice were fed a glucose meal and killed at different times later. Figure 4A (top) shows that serum glucose increased rapidly after glucose feeding, peaking at 10 min and then returning to a level twofold above fasted levels 50 min later. Serum insulin also increased quickly, but more transiently (Fig. 4B, middle). Insulin-stimulated expression of six insulin responsive loci was then examined (Fig. 4B and data not shown) including glucokinase (Gck) (Fig. 4A), an insulin-responsive gene whose product is involved in glycolysis (35,36). Gck mRNA was most robustly induced, with transcript levels increasing ninefold 60 min after insulin treatment (Fig. 4B, bottom). egr-1 had a large increase as well; however, this was mostly due to high levels of transcript in the livers of two mice, one at 20 and one at 30 min of glucose feeding (data not shown), while the transcript in most mice remained unchanged. Although the traditional view of IR signaling describes two divergent pathways, the mitogenic one transduced by B-Raf/MEK/Erk and the metabolic one transduced by PI3K/Akt (22), it has become increasingly apparent that MAPK pathways can also regulate metabolic state (16). Gck was chosen to examine recruitment of Pol II and signaling proteins because it had the highest, most consistent increase in transcript levels.

To show that the increase in Gck transcript was due to increased transcription, we used Matrix ChiP to determine Pol II levels along the locus. Gck has two transcription start sites (TSSs), 34,957 bp apart, with the downstream site being the one used by cells in the liver (the length from the start of the second TSS to the 3’ end is 14,301 bp) (35) (Fig. 4A). As expected, Pol II levels increased at the second TSS and the 3’ end of the gene (Fig. 4C, row 1). The 30-min peak at the 3’ end of the gene coincided with the first increase in Gck mRNA, suggesting that increased transcription was at least partially responsible for Gck induction (compare Fig. 4C, row 1, frame 3 with Fig. 4B, bottom).

Next we looked at the recruitment of IR complex components to Gck using Matrix ChiP. IRβ, Grb2, and SOS were recruited to the second TSS and 3’ end of Gck (Fig. 4C, rows 2, 4, and 5). Importantly, the phosphorylated, active form of IRβ was more robustly induced than the total receptor (Fig. 4C, row 3). These results are consistent with those seen in cell culture Figures 2 and 3.

Next we used Matrix ChiP to probe the recruitment of B-Raf, MEK, and ERK and their activated forms at the Gck locus in liver chromatin. Similar to the results in cells, the phosphorylated forms of all three kinases showed robust increases at all of the transcribed regions of the gene with kinetics similar to that of the phosphorylated insulin receptor (Fig. 4C, rows 7, 9, and 11). Also, as with the results in cells, ChiP with antibodies to the total kinases showed smaller responses compared with phosphorylated kinases (Fig. 4C, rows 6, 8, and 10). We also found a transient increase in the levels of MKP-1 at the second TSS and 3’ end of Gck. These data confirm that the insulin-induced recruitment and activation of the MAPK pathway seen in hepatocyte culture also occurs during insulin-induced expression of Gck in mammalian tissue.

Next, we compared Pol II and the phosphorylated kinases at first TSS and second TSS of Gck to the profiles at the TSS of gapdh, which, while highly expressed, was not stimulated in response to feeding within the 60-min time course (supplementary Fig. 3 [available in an online appendix], first row, compare frame 3 to frames 1 and 2). Interestingly, the constitutive Grb2, SOS, pB-Raf, pMEK, and pERK but not the pIRβ levels were much higher at gapdh than at the Gck. Also, unlike the feeding-induced increases in Grb2, SOS, and kinases seen at Gck, no such responses were seen at gapdh (supplementary Fig. 3). This result suggests that the glucose feeding-induced recruitment/activation of these MAPK signaling proteins at Gck but not gapdh is tightly coupled to the activated insulin receptor. Moreover, the fact that a similar type of inducible recruitment occurs in two different systems, one in vitro along a gene mediating mitogenic responses (Figs. 1–3) and another one in vivo along a gene involved in metabolism (Fig. 4), suggests that recruitment of IR/MAPK pathway in a pattern resembling Pol II elongation could be a general phenomenon of insulin-inducible genes.

**Absence of feeding-induced Gck mRNA response in ob/ob mice is associated with unsynchronized induction of IR-MAPK components at the Gck locus.** Obese (ob/ob) C57BL/6J mice are insulin resistant and have higher nuclear insulin receptor levels in the liver (13). Moreover, unlike the lean C57BL/6J mice, the ob/ob mice express constitutively high hepatic Gck activity that is unresponsive to either fasting or feeding (37,38). Thus, comparison of the lean and obese strains provides an opportunity to explore the relationship between direct recruitment of IR and MAPK components and Gck gene expression. We carried out side-by-side fasting followed by glucose feeding and performed the same measurements as in Fig. 4 in both lean and obese mice (Figs. 5–6). Compared with fasting lean mice, fasting ob/ob mice had
twice the serum glucose levels and >10-fold higher insulin levels (Fig. 5A). The fasting Gck mRNA and Pol II levels (Fig. 5A bottom and 5B, row 1) were also higher in the livers of the ob/ob mice, suggesting higher rates of transcription in the obese compared with the lean mice (Figs. 5A and B). The constitutive differences in Pol II levels at Gck gene (Fig. 5B row 1) in lean and obese mice matched the levels of IR (rows 2 and 3) and MAPK components (rows 4–6), suggesting that the baseline transcription and IR-MAPK activation is coupled in both strains. Histone H3
FIG. 5. Fasted obese mice have higher glucose, insulin, Gck expression, and Pol II and IR pathway recruitment than lean mice. A: Blood and serum from fasted lean and ob/ob mice were used to determine glucose and insulin concentrations, respectively (top and middle). Total RNA from livers of fasted lean and ob/ob mice was used in RT reactions with random hexamers. cDNA was used in PCR with primers to Gck and ribosomal protein L32 gene as a control for total RNA. B: Chromatin from the livers of fasted lean and ob/ob mice was used in ChIP assays with antibodies to the indicated proteins and histone H3 covalent modification. ChIPed DNA was used in PCR with primers to the indicated regions. C: Comparison of RT-PCR and ChIP results for Gck of fasted and 60-min glucose-fed lean and obese mice. Data (A–C) represent mean ± SEM of four mice for each time point.
lysine 9 acetylation (H3K9/14Ac), a mark of open chromatin, is typically high at the TSS and low at the 3' end of actively transcribed genes (39). Interestingly, at the Gck TSS, H3K9/14Ac but not H3 lysine 79 dimethylation (H3K79m2) (Fig. 5B rows 7 and 8) (40), a mark associated with transcription elongation, was higher in the ob/ob strain.
mice. Taken together, these results suggest that, at baseline, in the ob/ob mice, the hepatic chromatin structure at Gck is different from the lean mice and consistent with the higher transcription rates.

After glucose feeding, the transient increase in serum glucose was greater in the ob/ob mice but glycemic differences were not nearly as dramatic as the increases in insulin levels, which were >25-fold higher in the obese compared with the lean mice (Figs. 4 and 6A, middle). Unlike in the lean mice, the ob/ob mRNA levels did not respond to glucose feeding (Fig. 6A, bottom). Interestingly, after glucose feeding the Gck mRNA levels in the lean mice reached the unresponsive, constitutively high levels in the obese mice. The absent mRNA response is in agreement with the lack of induction in the 3′ end Pol II levels (Fig. 5C, row 2), where unlike the lean mice, little or no activation of IR and MAPK components was detected in the ob/ob strain (Fig. 5C, row 3, and 6B, rows 2–6).

Remarkably, glucose feeding in ob/ob mice induced second TSS Pol II, IR, and MAPK changes, but unlike the lean mice, these changes appeared unsynchronized and did not propagate to the 3′ end of the gene (Fig. 6B, rows 1–6). These results show that, unlike the lean mice, in the obese mice activation of Mek/Erk at Gck TSS is uncoupled from its canonical activator, B-Raf.

In contrast to the IR/MAPK changes (Fig. 6B, rows 1–6), the second TSS H3K9/14Ac levels in both lean and obese mice were largely unaffected by glucose feeding (Fig. 6B, row 7), indicating that this mark of open chromatin remains stable even though Pol II, IR, and MAPK activities are changing.

DISCUSSION

Here, using two inducible genes as examples, egr-1 in rat hepatocyte cultures and mouse Gck in liver tissues, we provide evidence that components of the IR signaling complex and downstream ERK MAPK cascade are directly recruited to insulin inducible loci. Our results suggest that effective mRNA synthesis/processing depends on coordinated activation of IR/MAPK cascade along Pol II transcribed regions.

Nuclear translocation of IR. Nearly 25 years ago, Olefsky and coworkers demonstrated nuclear translocation of IR and suggested that it may interact with specific targets in the genome to mediate insulin’s long-term effects on transcription (14). Our ChIP-based observations are consistent with this and other previous IR nuclear translocation studies (14,15). Nuclear translocation of insulin has also been demonstrated. Interestingly, insulin’s transcript through the nuclear pore may be independent of IR and instead be mediated by other cytoplasmic proteins (41). Using three different types of insulin antibodies, we could not detect a ChIP signal specific to insulin at either egr-1 or Gck (data not shown). Thus, it is conceivable that a large fraction of chromatin-bound IR is insulin-free.

Recruitment of IR and MAPK components to inducible gene loci. The IR/MAPK kinetics that resemble the Pol II pattern (Figs. 2–4, 6) suggest that these kinases either dictate or follow the Pol II initiation/elongation at the TSS. This raises the following question: what are the functional implications for this synchrony? Because many factors involved in chromatin rearrangements, gene transcription, and RNA processing are regulated by phosphorylation, coordinated action of signaling kinases at discrete points of time and space likely plays a critical role in synchronizing these events. The distribution of insulin-responsive kinases in the nuclear milieu could be too diffuse for the spatiotemporal specificity required to coordinate these processes (42,43). Thus, precisely timed recruitment and activation of kinases at induced gene loci could provide the requisite spatiotemporal specificity. For example, activated IR could serve to bring together and activate downstream components at target genes in a spatiotemporal manner that matches Pol II elongation. This tight spatiotemporal coupling would serve to compartmentalize kinase signals to specifically regulate productive Pol II mRNA synthesis from insulin-responsive gene loci. The side-by-side comparison of Gck responses in glucose-fed lean and obese mice lends further support to this suggestion (Fig. 6).

Differences and similarities of IR/MAPK recruitment to the Gck locus in lean and obese mice. Gck is a critical regulator of glucose metabolism, including hepatic control of glycogen synthesis and glycolysis (44). Thus, it would not be surprising if a chronic hyperglycemic state in the ob/ob mice were to trigger a homeostatic mechanism that maintains elevated constitutive Gck transcription to meet the metabolic burden posed by the high glucose load (Fig. 5A). The higher Gck mRNA levels in ob/ob mice, at least in part, reflect increased transcription as evidenced by higher Pol II density (Fig. 5B, row 1). The higher rates of transcription are further supported by the more open chromatin structure revealed by H3K9/14Ac differences between the two strains (Fig. 5B, row 7). The constitutive IR and MAPK ChIP signals at Gck in the lean and obese mice seem to match Pol II levels (row 1), suggesting that, in both strains, fasting activities of Pol II, IR (rows 1–3), and MAPK components (rows 4–6) are functionally coupled. After glucose feeding, Gck mRNA and 3′ end Pol II, IR, and MAPK densities in ob/ob mice remained unchanged. Interestingly, in lean mice, the mRNA and 3′ end values reached those of obese mice (Fig. 5C). This observation lends further support to the suggestion that 3′ end IR/MAPK cascade activities, coupled to Pol II, tune the mRNA output from the Gck locus to match the glucose load (Fig. 6A).

In sharp contrast to the strong similarity between IR/MAPK activities and Pol II density at the 3′ end during fasting in lean and obese mice (Fig. 5B), the changes in the levels of these proteins at the Gck TSS during feeding were dramatically different between the two strains (Figs. 4C and 6B). Specifically, the synchronous, feeding-induced increase in TSS pIR/B-Raf/MEK/ERK activities and Pol II in the lean mice (Figs. 4C and 6B) was absent in the ob/ob strain (Fig. 6B). Instead, in the obese mice at the TSS, there was a wave of Pol II, IR, and B-Raf activities, a pattern different from the steady MEK and Erk increase. The discrepant TSS kinase patterns in the obese mice suggest that at least a fraction of the total pool of MEK/Erk (rows 5 and 6) is uncoupled from the upstream activator, B-Raf, at this site (row 4). The apparent B-Raf–MEK/Erk uncoupling could reflect heterogeneous hepatocyte populations in the liver where activation of the MEK/Erk axis could be mediated by alternative upstream MEK activators such as C-Raf, A-Raf, or PAK (45,46,47).

Release of proximally paused Pol II, seen as a decrease in ChIP signal at the TSS, is an important mode of transcriptional activation (48). Whereas the initial feeding-induced decline (10 min) in TSS Pol II density in the ob/ob mice is consistent with this model (Fig. 6B, row 1), the increase that follows (20 min) may represent de novo
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promoter initiation. Remarkably, the TSS Pol II wave does not elongate to the 3' end; instead it seems that the feeding-induced polymerase either falls off or is stalled downstream. Thus, the unsynchronized activities of IR and MEK/Erk at the Gck TSS in ob/ob mice may be associated with low processivity (i.e., premature termination of RNA synthesis) (49) of the feeding-induced Pol II wave. Taken together, these observations are consistent with the suggestion that coupled activation of IR/B-Raf/MEK/Erk along a gene could play a role in productive mRNA synthesis by Pol II. In summary, we have shown for the first time that the IR signaling complex and downstream ERK MAPK cascade components are recruited and activated along insulin-induced loci. These observations raise many questions about how IR and the MAPK pathway are targeted to insulin-responsive genes and the role of this recruitment. Thus, it is clear that this facet of insulin- and IR-regulated gene expression needs to be explored. In this regard, the present study represents a new paradigm to study these processes at the chromatin level in health and disease.

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J.D.N. designed and performed experiments, analyzed the data, and wrote the manuscript. R.C.L. designed the experiments and wrote the manuscript. K.B. came up with the hypothesis, designed and performed experiments, analyzed the data, and wrote the manuscript.

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