The insulin-like growth factor 1 receptor (IGF1R) is a receptor tyrosine kinase with critical roles in various biological processes. Recent results from clinical trials targeting IGF1R indicate that IGF1R signaling pathways are more complex than previously thought. Moreover, it has become increasingly clear that the function of many proteins can be understood only in the context of a network of interactions. To that end, we sought to profile IGF1R–protein interactions with the proximity-labeling technique BioID. We applied BioID by generating a HEK293A cell line that stably expressed the BirA* biotin ligase fused to the IGF1R. Following stimulation by IGF1, biotinylated proteins were analyzed by MS. This screen identified both known and previously unknown interactors of IGF1R. One of the novel interactors was sorting nexin 6 (SNX6), a protein that forms part of the retromer complex, which is involved in intracellular protein sorting. Using co-immunoprecipitation, we confirmed that IGF1R and SNX6 physically interact. SNX6 knockdown resulted in a dramatic diminution of IGF1-mediated ERK1/2 phosphorylation, but did not affect IGF1R internalization. Bioluminescence resonance energy transfer experiments indicated that the SNX6 knockdown perturbed the association between IGF1R and the key adaptor proteins insulin receptor substrate 1 (IRS1) and SHC adaptor protein 1 (SHC1). Intriguingly, even in the absence of stimuli, SNX6 overexpression significantly increased Akt phosphorylation. Our study confirms the utility of proximity-labeling methods, such as BioID, to screen for interactors of cell-surface receptors and has uncovered a role of one of these interactors, SNX6, in the IGF1R signaling cascade.
we have discovered that the protein SNX6 plays an important role in regulating IGF1R signaling.

Results

BioID for the identification of novel IGF1R interactors

Our aim was to identify novel IGF1R interactors to further understand IGF1R signaling. To that end, we employed the proximity-dependent labeling technique BioID. We first generated a stable cell line that constitutively expressed the IGF1R as a fusion with BirA* (IGF1R–BirA*). To validate our stable cell line, we first performed immunofluorescence. IGF1R–BirA* was expressed largely at the plasma membrane, as is the case for native IGF1R (Fig. 1A). In contrast, cells expressing the BirA* construct showed uniform expression across the cell (Fig. 1A). Immunostaining also indicated robust biotinylation of vicinal proteins in the presence of biotin indicating that the BirA* within the IGF1–BirA* fusion was functional (Fig. 1B). Immunoblotting confirmed expression of the fusion construct at the expected molecular weight as well as efficient biotinylation of a number of proteins (Fig. 1C).

Following validation, we used the IGF1R–BirA* cell line to identify novel interactors of the IGF1R. This cell line was also used for all subsequent experiments.

These cells were exposed to vehicle or 100 nM IGF1, as well as an excess amount of biotin. Following treatment for 16 h, cells were lysed and streptavidin-conjugated beads were used to isolate biotinylated proteins. The identity of the biotinylated proteins was determined by LC-MS/MS. As shown in Table 1 we identified many of the previously known interactors of the IGF1R (see supporting Table 1 for additional detail). Using the STRING database, we generated an interaction network. Underscoring the power of the proximity-dependent labeling approach to identify protein interaction networks we identified not only known direct interactors of IGF1R but also proteins that interact with these interactors (Fig. 2).

The BioID screen identified a number of novel IGF1R interactors (STX5, VAPA, EPN4, RFIP5, RFIP1, and SNX6) that are involved in the movement of proteins between various cellular compartments. We chose to focus on sorting nexin 6 (SNX6), a key component of the retromer complex (25), as the mechanisms that control IGF1R trafficking, and its effect on IGF1R signaling, are unclear.

Confirmation of physical association between IGF1R and SNX6

To confirm the physical interaction between activated IGF1R and SNX6, we performed co-immunoprecipitation on
cells treated with IGF1, using an anti-IGF1R β antibody for pull-down. A distinct band corresponding to SNX6 was observed (Fig. 3), confirming association between this protein and the receptor.

**SNX6 knockdown perturbs IGF1R signaling but does not interfere with recycling dynamics**

To evaluate the contribution of SNX6 to IGF1R signaling, we initially performed siRNA-mediated knockdown of SNX6. Knockdown of SNX6 was robust (Fig. 4A). Importantly, SNX6 knockdown resulted in a significant diminution of IGF1-induced ERK1/2 phosphorylation 15 and 30 min following the addition of the ligand (Fig. 4, B–E). A similar trend was also noted for later time points but did not reach significance. Intriguingly, SNX6 knockdown did not result in a similar decrease in Akt phosphorylation.

We then assessed the effects of SNX6 knockdown on IGF1R internalization/recycling dynamics following IGF1 stimulation. A steady decrease in the relative amount of cell-surface IGF1 was observed following IGF1 treatment, as expected. SNX6 knockdown did not have a significant effect on the level of IGF1R internalization (Fig. 5A). In agreement with these results, there was no observable difference in the distribution of IGF1R between control and SNX6 knockdown cells (Fig. 5B).

**SNX6 knockdown suppresses recruitment of IRS1 and SHC1 to IGF1R**

To understand how SNX6 knockdown perturbs IGF1R signaling, we performed BRET assays to quantify the level of recruitment of the key signaling proteins, IRS1 and SHC1, to IGF1R in control and SNX6 knockdown cells. A clear suppression of IGF1-induced association of IRS1 and IRS1 or SHC1 can be seen in the case of SNX6 knockdown compared with control cells (Fig. 6).

**SNX6 overexpression promotes Akt phosphorylation in the absence of IGF1R stimulation**

We next decided to assess the effects of increasing SNX6 protein levels on IGF1R signaling. Over-expression of SNX6 had no significant effect on IGF1-stimulated ERK1/2 or Akt phosphorylation (Fig. 7, A and B). Of note, SNX6 over-expression resulted in a pronounced increase in Akt phosphorylation even in the absence of any ligand. This level of Akt phosphorylation was comparable with the level induced by IGF1 itself (Fig. 7, A and B). A similar effect on ERK1/2 phosphorylation was not observed (Fig. 7, A and B).

**SNX6 overexpression has no effect on IGF1R distribution in the cell**

Finally, we chose to assess the effects of increased SNX6 protein levels on the cellular distribution of IGF1R. We observed no obvious effects of increased SNX6 protein levels on the distribution of IGF1R in the basal state or following IGF1 stimulation (Fig. 7C).

**Discussion**

Recent advances in proteomics have enabled a finer dissection of both the insulin and IGF1 signaling pathways. For example, quantitative phosphoproteomic analysis has been used to identify novel participants in the insulin (and likely IGF1) signaling pathway (26). This type of analysis has even been used to quantify changes in the phosphorylation of IGF1R itself (27). Temporal proteomic analysis has revealed that IGF1 treatment over time promotes the production of proteins associated with vesicle trafficking, glycolysis, and cell stress, functions that have been associated with cancer cell proliferation (28). In addition to being used to analyze cell signaling events, proteomics has also been used to identify proteins that interact with the IGF1R and its adaptors (29). Classical approaches for the identification of protein interactions have several limitations that preclude identification of many biologically important interactions. This barrier has been overcome by proximity-dependent labeling methodologies. To enable a more comprehensive characterization of the IGF1R interactome, one that includes weak, indirect, or transient interactors, we made use of the proximity-labeling method known as BioID.

BioID has been extensively used to define interaction networks for a number of proteins and subcellular regions (reviewed in Ref. 30). APEX is an example of another popular proximity-labeling method that has been used for similar studies (16). To our knowledge there have been only two published studies that have used this approach to define theTable 1

| Primary Protein Name | Protein Description | Fold IGF1 vs NT | t-test IGF1 vs NT |
|----------------------|----------------------|----------------|------------------|
| PTN11                | Tyrosine-protein phosphatase non-receptor type 11 | 19.6 ± 0.000 | 0.000 |
| SHC1                 | SHC-kinase activating protein 1 | 17.5 ± 0.004 | 0.000 |
| M4K5                 | Mitogen-activated protein kinase kinase kinase kinase | 6.9 ± 0.002 | 0.000 |
| SWP70                | Switch-associated protein 70 | 5.1 ± 0.001 | 0.000 |
| DDRGK                | DDRGK domain-containing protein 1 | 3.3 ± 0.007 | 0.000 |
| ZFY16                | Zinc finger FYVE domain-containing protein 16 | 3.2 ± 0.025 | 0.000 |
| RFP5                 | Rab11 family-interacting protein 5 | 3.1 ± 0.018 | 0.000 |
| MPRA                 | Canonical mannose-6-phosphate receptor | 3.1 ± 0.008 | 0.000 |
| TOR1A                | Torin-1A-interacting protein 1 | 3.0 ± 0.000 | 0.000 |
| SHIP2                | Phosphotyrosinolysol-3,4,5-trisphosphate 5-phosphatase 2 | 2.8 ± 0.002 | 0.000 |
| GRP78                | Glucose-regulated protein 78 | 2.7 ± 0.001 | 0.000 |
| MAF4                 | Microtubule-associated protein 4 | 2.6 ± 0.011 | 0.000 |
| UBE2J                | Ubiquitin-conjugating enzyme E2 J1 | 2.6 ± 0.000 | 0.000 |
| NCK1                 | NCK1 cytoplasmic protein | 2.6 ± 0.001 | 0.000 |
| STX5                 | Syntaxin-5 | 2.5 ± 0.003 | 0.000 |
| RFP1                 | Rab11 family-interacting protein 1 | 2.4 ± 0.000 | 0.000 |
| RRR5                 | Retin receptor | 2.3 ± 0.002 | 0.000 |
| CNP3                 | CANP-3 | 2.3 ± 0.008 | 0.000 |
| LRC50                | Leucine-rich repeat-containing protein 59 | 2.3 ± 0.001 | 0.000 |
| CDKAL                | Threonine/myristoyl-ladenosine RNA methyltransferase | 2.2 ± 0.006 | 0.000 |
| SRPRA                | Signal recognition particle receptor subunit alpha | 2.2 ± 0.014 | 0.000 |
| PTN11                | Tyrosine-protein phosphatase non-receptor type 11 | 2.2 ± 0.005 | 0.000 |
| VAPA                 | Vesicle-associated membrane protein-activated protein A | 2.2 ± 0.002 | 0.000 |
| KTH41                | Kinesin | 2.2 ± 0.015 | 0.000 |
| EPN4                 | Epsin-4 | 2.2 ± 0.002 | 0.000 |
| RTN4                 | Retinol-4 | 2.1 ± 0.001 | 0.000 |
| LYRIC                | Protein LYRIC | 2.1 ± 0.003 | 0.000 |
| BAP31                | B-cell receptor-associated protein 31 | 2.1 ± 0.006 | 0.000 |
| RBP4P1               | Ribo-3' binding protein 1 | 2.1 ± 0.025 | 0.000 |
| STIM1                | Stromal interaction molecule 1 | 2.0 ± 0.033 | 0.000 |
| SNX6                 | Sorting nexin-6 | 2.0 ± 0.002 | 0.000 |
| KAD9                 | Adenylyl kinase 9 | 2.0 ± 0.026 | 0.000 |

**SNX6 is a novel regulator of IGF1R signaling**

To understand how SNX6 knockdown perturbs IGF1R signaling, we performed BRET assays to quantify the level of recruitment of the key signaling proteins, IRS1 and SHC1, to IGF1R in control and SNX6 knockdown cells. A clear suppression of IGF1-induced association of IRS1 and IRS1 or SHC1 can be seen in the case of SNX6 knockdown compared with control cells (Fig. 6).
interactome of cell-surface receptors (31, 32), and in both cases G protein–coupled receptors were the subject of these studies. In this paper we describe the first use of a proximity-labeling method to characterize the interactome of a receptor tyrosine kinase.

Our screen yielded a number of well-established interactors of the activated IGF1R, such SHC1 and PTPN11. These findings underscore the validity of the BioID technique to define the complete IGF1R interactome. One of the novel candidates we identified was SNX6. SNX proteins have been shown to play roles in the recycling and signaling events of a number of cell-surface receptors (33). There are very few published studies on the role of SNX proteins in RTK signaling and function. However, there is one study that links SNX6 with the insulin receptor, a receptor that is very similar to IGF1R (34). Given this information, we chose to further examine the role of SNX6 in IGF1R signaling and function.

We confirmed physical interaction between IGF1R and SNX6 by co-immunoprecipitation. We then showed that SNX6 knockdown resulted in a diminution of IGF1-induced ERK1/2 phosphorylation. A possible explanation for this observation could be that SNX6 knockdown interferes with IGF1-induced IGF1R internalization, which is necessary for the activation of the ERK1/2 pathway (35). However, we found that the internalization dynamics were not significantly perturbed by SNX6 knockdown.

![Figure 2. Known and predicted interactions between the proteins in Table 1 according to the STRING database.](image)

![Figure 3. Co-immunoprecipitation confirms physical interaction of IGF1R and SNX6. Pulldown was performed with either isotype control or anti-IGF1Rβ antibody, followed by immunoblot with an anti-SNX6 antibody or an anti-IGF1Rβ antibody, as indicated. A distinct band corresponding to SNX6 can be seen for the IGF1Rβ pulldown lane. This blot is representative of three independent experiments.](image)
To achieve a mechanistic understanding of how SNX6 knockdown perturbs IGF1R signaling, we performed BRET to quantify recruitment of the key signaling adaptor proteins, IRS1 and SHC1, to IGF1R. We found that knocking down SNX6 inhibited the association between IGF1R and both IRS1 and SHC1. Although this would explain why IGF1-induced ERK1/2 phosphorylation is diminished in SNX6 knockdown cells, as both IRS1 and SHC1 recruitment promote downstream ERK1/2 phosphorylation, it does not explain why we did not see any effect on Akt phosphorylation. Perhaps Akt phosphorylation requires only a modest level of IGF1R activation. Alternatively, there might be compensation by the closely-related IRS2 isoform, as these two proteins have both been shown to promote Akt phosphorylation (36).

Figure 4. SNX6 knockdown impairs IGF1R signaling. A, a representative blot showing robust siRNA-mediated knockdown of SNX6. B, representative blot showing that SNX6 knockdown inhibited IGF1-induced phosphorylation of ERK1/2, whereas having no significant effect on IGF1-induced Akt phosphorylation. C, quantification of bands in B. D, a representative blot highlighting a similar trend for later treatment times. E, quantification of bands in D. One-way analysis of variance was performed followed by Bonferroni’s Multiple Comparison Test, 0.0001 < ****, p < 0.001; ***, p < 0.001; *, p < 0.05. n = 3–4.
Conversely, SNX6 overexpression had no significant effect on either IGF1-induced ERK1/2 or Akt phosphorylation. Intriguingly, SNX6 overexpression alone resulted in a significant increase in Akt phosphorylation, which was comparable with the level of phosphorylation induced by IGF1.

A recent study showed that blocking IGF1R internalization inhibited both Akt and ERK1/2 phosphorylation; leading the authors to conclude that much of IGF1R signaling takes place at the endosome (37). SNX6 is associated with the early endosome (25) presumably by binding of its PX domain to phosphatidyli-
Nonsitol 3-phosphate, which is the most abundant phosphoinositide in the early endosome membrane (38). Both Akt (39) and ERK1/2 (40) have also been shown to signal from the early endosome. Therefore, SNX6 might promote the association of IGF1R with the early endosomal compartment where it will be in close proximity to members of its signaling cascade, including Akt and ERK1/2. This might also serve to explain why we saw an increase in Akt as a result of SNX6 overexpression even in the absence of IGF1 treatment. Perhaps SNX6 overexpression enhances the association of IGF1R with the early endosome. This would serve to (a) bring IGF1R in close proximity to its substrate proteins and (b) draw it away from the degradative lysosomal pathway. Therefore, the effects of even low levels of basal IGF1R kinase activity would be amplified. To test these hypotheses, it would be instructive to measure the effects of SNX6 overexpression on the occupancy (both in terms of quantity and time spent) of IGF1R, Akt, and ERK1/2 in the early endosome.

In summary, the data presented here confirm the utility of using proximity-labeling methods, such as BioID, to identify
SNX6 is a novel regulator of IGF1R signaling

novel interactors of a cell-surface receptor. Importantly, we have validated the role of SNX6 as a novel mediator of IGF1R function; revealing a hitherto unknown link between SNX/retromer and IGF1R signaling. In addition to its biological significance, this finding also presents an additional target for the therapeutic modulation of IGF1R signaling.

Experimental procedures

BioID principle

BioID was originally developed by Dr. Kyle Roux (14) and makes use of a promiscuous biotin ligase known as BirA*; a mutated version of the BirA ligase. BirA catalyzes a two-step reaction in the presence of biotin. In the first step, BirA combines biotin and AMP to produce biotinoyl-5’-AMP (bioAMP); an activated form of biotin. This activated biotin is retained by BirA in its active site until it comes into contact with a specific lysine residue in a minimal recognition sequence known as the biotin acceptor tag. BirA* also produces the activated biotin bioAMP. However, BirA* does not retain bioAMP in its active site. Instead, BirA* prematurely releases the bioAMP; where it reacts with primary amines of adjacent proteins. This property of BirA* is harnessed in the BioID technique by fusing it with a protein of interest.

BioID construct and generation of the IGF1R–BirA* stable cell line

To generate the IGF1R–BirA* construct, the coding region of human IGF1R (IGF1R; GenBank accession no. NM_000875) minus the stop codon was subcloned into the pcDNA3.1 MCS-BirA(R118G)-HA plasmid (Addgene) such that BirA* was expressed on the C-terminal (cytoplasmic) end of the IGF1R. A stable cell line was generated by stably transfecting HEK293A cells with the IGF1R–BirA* fusion construct, which is referred to as the “BioID cell line” for the rest of this paper. The IGF1R–BirA* construct was transfected into the HEK293A cells with the Lipofectamine LTX transfection reagent (Thermo Fisher Scientific). As this construct contains a neomycin cassette, stable transfectants were selected for by culturing the cells in media containing 1 mg/ml of the antibiotic G418 (Life Technologies). The BioID cell line was subsequently cultured in normal HEK293A growth media supplemented with 0.5 mg/ml of G418.

Quantitative MS

Sample preparation—Following treatment, cells were lysed and pulldown was performed using streptavidin-coated beads according to a previously published BioID protocol (41). Samples were submitted to the Duke Proteomics Core Facility (DPCF) in 30 μl of 250 mM Tris, 3% SDS. The entire sample was loaded onto an Invitrogen NuPAGE 4–12% SDS-PAGE gel and run for ~5 min to electrophorese all proteins into the gel matrix. The entire Mț range was then excised in a single gel-band and subjected to standardized in-gel reduction, alkylation, and tryptic digestion. Following lyophilization of the extracted peptide mixtures, samples were resuspended in 20 μl of 2% acetonitrile, 1% TFA supplemented with 10 fmol/μl of yeast alcohol dehydrogenase. From each sample, 5 μl was removed to create a QC Pool sample that was run periodically throughout the acquisition period.

Quantitative analysis method—Quantitative LC/MS/MS was performed on 2 μl of each sample, using a nanoAcquity UPLC system (Waters Corp) coupled to a Thermo QExactive Plus high resolution accurate mass tandem mass spectrometer (Thermo) via a nanoelectrospray ionization source. Briefly, the sample was first trapped on a Symmetry C18 20-mm × 180-μm trapping column (5 μl/min at 99.9/0.1 (v/v) water/acetonitrile), after which the analytical separation was performed using a 1.7-μm Acquity BEH130 C18 75-μm × 250-mm column (Waters Corp.) with a 90-min linear gradient of 5–40% acetonitrile with 0.1% formic acid at a flow rate of 400 nanoliters/min with a column temperature of 55 °C. Data collection on the QExactive Plus mass spectrometer was performed in a data-dependent acquisition mode of acquisition with a r = 70,000 (m/z 200) full MS scan from m/z 375–1600 with a target AGC value of 1e6 ions followed by 10 MS/MS scans at r-17,500 (m/z 200) at a target AGC value of 5e4 ions. A 20-s dynamic exclusion was employed to increase depth of coverage. The total analysis cycle time for each sample injection was ~2 h.

Following UPLC-MS/MS analyses, data were imported into Rosetta Elucidator version 4.0 (Rosetta Biosoftware, Inc.), and analyses were aligned based on the accurate mass and retention time of detected ions (features) using the PeakTeller algorithm in Elucidator. Relative peptide abundance was calculated based on area under the curve of the selected ion chromatograms of the aligned features across all runs. The MS/MS data were searched against a custom Swissprot database with Homo sapiens taxonomy (downloaded in Sept. 2016) with additional proteins, including yeast ADH1, BSA, streptavidin, Escherichia coli BirA as well as an equal number of reversed-sequence (“decys”) false discovery rate determination. Mascot Distiller and Mascot Server (version 2.5, Matrix Sciences) were utilized to produce fragment ion spectra and to perform the database searches. Database search parameters included fixed modifications on Cys (carbamidomethyl) and variable modifications on Asn and Gln deamidation. After individual peptide scoring using the PeptideProphet algorithm in Elucidator, the data were annotated at a 0.5% peptide false discovery rate.

Statistical analysis—Fold-changes for every identified protein was calculated between the IGF1-treated and untreated groups. In addition, we performed a two-tailed heteroscedastic t test in Excel for each of these comparisons. To select proteins that were differentially interacting with IGF1R, proteins were filtered to include those with a positive 2-fold change and p < 0.05 compared with untreated group. We required proteins to have a ProteinTeller score of >0.8 to maintain a high qualitative identification probability.

Knockdown and overexpression

ON-TARGET plus siRNA pools (Dharmacon) were used to suppress the expression of human SNX6. Transfection was performed 1 day after seeding using the Lipofectamine RNAiMAX reagent (Invitrogen) following the manufacturer’s instructions. A nontargeting siRNA pool was used as a negative control. SNX6 overexpression was achieved by transfecting cells with a plasmid containing the coding region of human SNX6 un-
under the influence of a cytomegalovirus promoter (catalog CH803743; Vigene Biosciences) using the Lipofectamine LTX transfection reagent (Thermo Fisher Scientific) according to the manufacturer’s instructions.

**Immunoblotting**

Protein extraction from cultured cells and Western blotting was performed according to a previously published protocol (12). Proteins were resolved on an SDS-PAGE (Life Technologies) and transferred to a nitrocellulose membrane (Bio-Rad). The following antibodies were used: phospho-Akt(S473) number 9271, Akt number 9272, phospho-ERK1/2 number 9101, ERK1/2 number 9102, DYKDDDDK Tag (9A3) number 8146 (all from Cell Signaling Technology), and monoclonal anti-SNX6 antibody (S6324; Sigma). All antibodies were applied at a 1:1000 concentration. Proteins were visualized by chemiluminescence using ECL Prime reagent (GE Healthcare) and a Syngene G:Box. Band intensities were determined using ImageJ software.

**BRET**

Constructs used—The SHC1(PTB)–eYFP and IRS1(PTB)–eYFP constructs were generated by subcloning the fragments encoding the phosphotyrosine binding (PTB) domain of human SHC1 and the PTB domain of human IRS1, respectively, into the pEYFP-N1 vector. The IGF1R–RlucII construct was generated by subcloning the coding region of the human IGF1R into the pcDNA3.1 RlucII vector (a kind gift from Thomas Pack, Duke University). All constructs were validated by sequencing.

Immunofluorescence

Immunofluorescence was performed by magnetic separation using protein A-agarose beads (number 73778) according to the manufacturer’s protocol (Cell Signaling Technology). Cells were seeded at a density of 500,000 cells per 10-cm plate. Pulldown was performed using either IGF1Rβ antibody 3027 or mouse anti-rabbit IgG (Light-Chain Specific) (D4W3E) mAb 45262 at a concentration of 1:100 using identical amounts of cell lysate (200 µg) (Cell Signaling Technology).

**Flow cytometry**

Following treatment, cell collection was performed in enzyme-free cell dissociation buffer (Life Technologies). Cells were fixed with either phycoerythrin-conjugated mouse anti-human CD221 antibody (catalog number 555999; BD Biosciences) or phycoerythrin-conjugated mouse IgG1 Isotype Control (12-4714-42; eBioscience) diluted at 1:5 in incubation buffer (0.5 g of BSA in 100 ml of 1× PBS) for 1 h at room temperature in the dark. Following two final washes in 1× PBS, flow cytometry was performed using a BD FACS Canto II flow cytometer (BD Biosciences) and data analysis was performed using FlowJo Version 10.

**Immunofluorescence**

Following treatment, cells were fixed with 4% paraformaldehyde for 15 min at room temperature. Following fixation, cells were washed twice with 1× PBS. 10 mM NH4Cl/PBS was added for 20 min to quench the free aldehyde groups of the fixative. Following three more washes with 1× PBS, cells were permeabilized in 0.1% Triton X-100 (in PBS) for 10 min. Following an additional three washes with 1× PBS, 5% normal goat serum was added for 10 min. Cells were then stained using an HA tag (C29F4) antibody (1:200; Cell Signaling Technology) overnight at 4 °C. The following day, cells were washed and blocked again in 5% normal goat serum. Alexa Fluor secondary antibodies (Invitrogen) were used for detection. Cells were counterstained with 4′,6-diamidino-2-phenylindole to visualize nuclei. Images were taken using an LSM 510 META confocal microscope (Zeiss) with ZEN software.

**Statistics**

For pairwise comparisons, data were analyzed using Student’s t test. For multiple comparisons, one-way analysis of variance followed by Bonferroni’s Multiple Comparison Test. All data are presented as mean ± S.E. A value of $p < 0.05$ was considered to be statistically significant. Statistical analysis was performed on GraphPad Prism 5.0.

**Author contributions**—A. B., C. P. H., and V. J. D. conceptualization; A. B., E. S., and G. W. data curation; A. B., C. P. H., and V. J. D. formal analysis; A. B., C. P. H., and V. J. D. investigation; A. B., E. S., and G. W. methodology; A. B., C. P. H., and V. J. D. writing-original draft; A. B., C. P. H., and V. J. D. writing-review and editing; C. P. H. and V. J. D. supervision; E. S. resources; V. J. D. funding acquisition.

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