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Critical evaluation of the expression of gastrin-releasing peptide in dorsal root ganglia and spinal cord

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Abstract
There are substantial disagreements about the expression of gastrin-releasing peptide (GRP) in sensory neurons and whether GRP antibody cross-reacts with substance P (SP). These concerns necessitate a critical revaluation of GRP expression using additional approaches. Here, we show that a widely used GRP antibody specifically recognizes GRP but not SP. In the spinal cord of mice lacking SP (Tac1 KO), the expression of not only GRP but also other peptides, notably neuropeptide Y (NPY), is significantly diminished. We detected Grp mRNA in dorsal root ganglia using reverse transcription polymerase chain reaction, in situ hybridization and RNA-seq. We demonstrated that Grp mRNA and protein are upregulated in dorsal root ganglia, but not in the spinal cord, of mice with chronic itch. Few GRP+ immunostaining signals were detected in spinal sections following dorsal rhizotomy and GRP+ cell bodies were not detected in dissociated dorsal horn neurons. Ultrastructural analysis further shows that substantially more GRPergic fibers form synaptic contacts with gastrin releasing peptide receptor-positive (GRPR+) neurons than SPergic fibers. Our comprehensive study demonstrates that a majority of GRPergic fibers are of primary afferent origin. A number of factors such as low copy number of Grp transcripts, small percentage of cells expressing Grp, and the use of an eGFP GENSAT transgenic as a surrogate for GRP protein have contributed to the controversy. Optimization of experimental procedures facilitates the specific detection of GRP expression in dorsal root ganglia neurons.

Keywords
substance P, gastrin-releasing peptide, itch, dorsal root ganglia, sensory neurons, spinal cord

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Introduction

Many earlier studies have demonstrated the specificity of gastrin-releasing peptide (GRP) antibodies by radioimmunoassay and immunohistochemistry (IHC) after dorsal rhizotomy or antibody absorption, and have concluded that GRP-positive (GRP+) fibers, or GRP content in the spinal cord, are predominantly of peripheral origin.1-5 In addition, it was shown that GRP antibody does not cross-react with substance P (SP), and that GRP is present in dorsal root ganglia (DRGs) of cat, dog, rat, and monkey.2-7

After more than a decade of relative dormancy, the identification of GRP as an itch-transmitting peptide in DRGs has reignited considerable interest in GRP expression.8-12 We propose that GRP is a key itch-specific neuropeptide in sensory neurons that is released from primary afferents to activate postsynaptic GRPR in the spinal cord in response to nonhistaminergic stimuli.12-16

Consistent with this notion as well as with many earlier studies, several other laboratories have independently shown that the GRP antibody recognizes a distinct subset of DRG neurons mostly using 1:1000 or 1:4000 dilution.17-20 Moreover, GRP is up-regulated in DRGs of mice and monkeys with chronic itch12,13,21,22 and a portion of GRP+ neurons are distinct from SP+ neurons in DRG.18,23 More recently, 3-D synaptome analysis using high-voltage electron microscopy and chemical anatomy revealed that the electron-dense GRP+ vesicles are present in presynaptic afferents contacting postsynaptic neurons with the electron-dense postsynaptic densities in the spinal cord.24

On the other hand, conflicting results on GRP expression in DRGs using the GRP antibody have also been reported (Table 1). Most notably, some researchers were unable to detect specific GRP immunostaining in DRGs. The issue was raised about the dilution of the antibody (Immunostar) used in the studies. Solorzano et al.25 showed that at 1:1000 dilution GRP immunostaining is widespread in DRGs, similar to several previous studies.26,27 On the other hand, using 1:1000 dilution, Fleming et al.28 showed only 1.79% of DRGs are positive for GRP, the smallest percentage ever reported. While Solorzano et al.25 argued that at 1:4000 dilution, GRP antibody is specific but could not detect any staining in DRG, Kiguchi et al.17 recently showed distinct staining in a small subset of DRGs using the same 1:4000 dilution.

Detection of Grp mRNA by in situ hybridization (ISH) in DRGs also remains controversial. Although we were able to observe Grp expression in DRGs by ISH,12,29 others could not detect positive signals.25,28,30,31 Moreover, two groups did not detect Grp mRNA in DRG by RNA-seq.32,33 While several laboratories detected Grp mRNA by reverse transcription polymerase chain reaction (RT-PCR) using single cell method,26,27 Solorzano et al.25 argued that the detections are due to de novo expression in DRG neuron culture conditions. On the other hand, it has been reported that Grp mRNA was detectable from uncultured DRGs by RT-PCR,28,30 qRT-PCR12 and a cDNA microarray study34 (Table 1).

Two recent studies argued that the widely used GRP antibody cross-reacts with SP25,33 because GRP immunostaining is reduced in mice lacking Tac1.25 In contrast, others showed specific GRP and SP double immunostaining in DRGs, arguing against this possibility.18,27

These discrepancies in the literature prompted us to revisit the issue, with a focus on potential reasons that may explain some of the inconsistent results and newly

| Method       | GRP expression                          | References |
|--------------|-----------------------------------------|------------|
| IHC/IF (WT)  | + (lab derived); + (1:1000); IS         | 2, 3, 4    |
|              | + (1:100); IS                            | 16, 20, 28 |
|              | +++ (1:1000); IS                         | 26, 29     |
|              | + (not indicated)                        | 19         |
|              | + (1:500); IS                            | 13, 23     |
|              | ++++ (1:100); SCBT                       | 27         |
|              | + (1:1000, 1:2000); Assaypro             | 18         |
|              | ++++ (1:1000), - (1:4000); IS            | 25         |
|              | + (1:4000); IS                           | 17         |
| IHC/IF (Grp KO) | - (1:1000); IS                        | 29         |
|              | - (1:500); IS                            | 13         |
|              | ++++ (1:1000), - (1:4000); IS            | 25         |
| ISH          | - (bases 212–634)                        | 28         |
|              | - (coding region, bases 115–554)        | 25         |
|              | - (not indicated)                        | 30, 31     |
|              | + (bases 149–707)                        | 12         |
| RT-PCR gel electrophoresis | + (weak band)                          | 28         |
|              | + (single cell)                          | 26, 27     |
| qRT-PCR      | + (trace amount)                        | 30         |
|              | + (low levels)                           | 25         |
| RNA-Seq      | + (not indicated)                       | 33         |
|              | - (single cell)                         | 32         |
| cDNA Microarray | +                                    | 34         |

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- not detected or absent, + detected and/or distinct staining pattern, +++ wide spread staining pattern; IHC/IF antibody dilutions are indicated in parentheses; ISH Grp mRNA region used for antisense probe indicated in parentheses (NCBI accession NM_175012.4), IS: Immunostar; SCBT: Santa Cruz Biotechnology.
Materials and methods

Animals

Male mice between 7 and 12 weeks old were used for experiments. C57BL/6 J mice were purchased from the Jackson Laboratory (http://jaxmice.jax.org/strain/013636.html). C57BL/6 J mice, GRPR-eGFP BAC Transgenic mice from MMRRC (i.d. 036178), Grp KO,13 TacI KO,35 BrafNaV1.8,13, and their respective wild type (WT) littermates were used. All mice were housed under a 12 h light/dark cycle with food and water provided ad libitum. All experiments were performed in accordance with the guidelines of the National Institutes of Health and the International Association for the Study of Pain and were approved by the Animal Studies Committee at Washington University School of Medicine.

Ablation of TRPV1⁺ fibers

C57BL/6 J mice were treated with resiniferatoxin (RTX) (25 ng in 5 μL, intrathecal) as previously described, with a modification in the dose of RTX.36 Seven days after RTX injection, mice were perfused, and lumbar spinal cord tissues were collected for immunostaining.

Dorsal rhizotomy

C57BL/6 J male mice were used for unilateral rhizotomy at spinal lumbar level L4–L6.13 Briefly, laminectomy was performed to expose the L4–L6 dorsal roots, which were sharply transected. Animals were perfused, and the lumbar spinal cord tissues were collected 14 days after the dorsal rhizotomy for immunostaining.

Xerosis (dry skin) model

The AEW (acetone-ether-water) dry skin model was implemented as described.37,38 Briefly, the nape of mice was shaved and a mixture of acetone and diethyl ether (1:1) was applied with a cotton pad on the neck skin for 15 s, followed immediately by a 30 s distilled water application. This regimen was administered twice daily for eight days. Littermate control mice received water only for 45 s on the same schedule. Spontaneous scratches were counted for 60 min on the morning following the last AEW treatment. On day 8, AEW-treated mice displayed 150–300 scratching bouts in 60 min. Cervical and thoracic DRG and spinal cord tissues were isolated and processed for IHC, ISH, and RT-PCR and qRT-PCR.

Immunohistochemistry

Mice were anesthetized (ketamine, 100 mg/kg and Xylazine, 15 mg/kg) and perfused intracardially with PBS pH 7.4 followed by 4% paraformaldehyde (PFA) in PBS. Tissues were dissected, post-fixed for 2–4 h, and cryoprotected in 20% sucrose in PBS overnight at 4°C. Tissues were sectioned in OCT using a cryostat microscope. IHC was performed as described.14 Briefly, free-floating frozen sections at 20 μm thickness were blocked in a 0.01 M PBS solution containing 2% donkey serum and 0.3% Triton X-100 followed by incubation with primary antibodies overnight at 4°C, washed three times with PBS, secondary antibodies for 2 h at room temperature, and washed again three times. For biotin-conjugated secondary antibodies, sections were next incubated with avidin-conjugated fluorophores and washed three times. Sections were mounted on slides and approximately 100 μL FluoromountG (Southern Biotech) was placed on the slide with a coverslip. Fluorescein isothiocyanate (FITC)-conjugated Isolectin B4 from Griffonia simplicifolia (IB4, 10 μg/mL; L2895, Sigma) or the following primary antibodies were used, rabbit anti-GRP (1:500–1:4000; Immunostar, 20073, lot #1420001), rabbit anti-calcitonin gene-related peptide alpha (CGRPα) (1:5000; Millipore, AB15360), guinea pig anti-CGRP,18 guinea pig anti-SP (1:1000; Abcam, ab10353, lot# GR29977-17), guinea pig anti-transient receptor potential cation channel subfamily V member 1 (TRPV1) (1:1000; Peninsula Labs, T-5027), guinea pig anti-SP (1:1000; Abcam, ab10353, lot# GR29977-17), guinea pig anti-transient receptor potential cation channel subfamily V member 1 (TRPV1) (1:1000; NeuroMics, GP14100), and chicken anti-GFP antibody (1:500; Aves Labs, GFP-1020). For GRP/GRP/SP triple staining, a total of 10 adult GRPR-eGFP male mice and chicken anti-GFP antibody (1:500; Aves Labs) were used. The secondary antibodies were FITC-, Cyanine 3 (Cy3)-, Cy5 donkey anti-guinea pig (1:500; Millipore) or Alexa 594 conjugated donkey anti-rabbit or anti-guinea pig IgG (1:500, Jackson ImmunoResearch), or biotin-SP-conjugated donkey anti-rabbit or anti-chicken IgG (1:400, Jackson ImmunoResearch) and Neutravidin-conjugated Alexa Fluor488 (1:1000, Life Technologies), Third antibody—FITC-avidin (1:1000; Vectorlabs). Fluorescent Images were taken using a Nikon Eclipse Ti-U microscope with CoolSnapHQ CCD Camera (Photometrics). Staining intensities for each section were quantified by an observer blinded to the group or
genotype using ImageJ (version 1.34e, NIH Image) as previously described.\textsuperscript{13}

**DRG and spinal dorsal horn neuron cultures**

Primary cultures of DRGs and spinal dorsal horn neurons were prepared from seven-weeks-old C57BL/6J mice.\textsuperscript{13} Mice were sacrificed, DRGs and dorsal horn of spinal cord were dissected out and incubated, separately, in Neurobasal-A Medium (Gibco) containing 10 μL papain (Worthington) at 37°C for 20 min, and an additional 20 min digestion at 37°C for DRGs with collagenase type 2 (Worthington). Enzymatic digestion was stopped by adding another 2 mL Neurobasal-A medium. After washing with the same medium for three times, gentle trituration was performed using flame polished glass pipette until the solution became cloudy. The homogenate was centrifuged at 500 g for 5 min, and the supernatant was discarded. Cell pellets were resuspended in culture medium composed of Neurobasal medium (Gibco, 92% vol/vol), fetal bovine serum (Invitrogen, 1% vol/vol), B27 (Invitrogen, 2% vol/vol), GlutaMax (2 mM, Invitrogen, 1% vol/vol), Penicillin (100 μg/mL) and Streptomycin (100 μg/mL) and then plated onto 12-mm coverslips coated with laminin and poly-ornithine for 1 h.

GRP immunostaining on dissociated DRG neurons and dorsal horn neurons was performed as previously described.\textsuperscript{39} Cells were fixed in 4% paraformaldehyde for 1 h at room temperature and blocked in blocking buffer (PBS containing 2% normal donkey serum and 0.3% Triton X-100) for 1 h at room temperature. Cells were then incubated with rabbit anti-GRP antibody (ImmunoStar, 1:1,000) and mouse anti-NeuN antibody (EMD Millipore, 1:1,000) in blocking solution at room temperature for 2–16 h for colorimetric detection. Reactions were stopped by washing in 0.5% paraformaldehyde in PBS. Bright field images were taken using a Nikon Eclipse Ti-U microscope with a Nikon DS-Fi2 Camera. ISH-positive and negative neurons were quantified by an observer blinded to the group or genotype using ImageJ (version 1.34e, NIH Image) as previously described.\textsuperscript{13}

**In situ hybridization**

ISH was performed as previously described,\textsuperscript{41} using a digoxigenin-labeled cRNA (Roche) antisense probe for Grp (bases 149-707 of Grp mRNA, NCBI accession NM_175012.4) and Tac1. Briefly, on-slide frozen cervical and thoracic DRG sections at 20 μm thickness were incubated with Proteinase K (50 μg/mL) buffer for 10 min, incubated in prehybridization solution for 3 h at 65°C and then incubated with Grp or Tac1 probe (~2 μg/mL) in hybridization solution overnight at 65°C. After SSC stringency washes and RNase A (0.1 μg/mL for 30 min) incubation, sections were incubated in 0.01 M PBS with 20% sheep serum and 0.1% Tween blocking solution for 3 h and then incubated with anti-digoxigenin antibody conjugated to alkaline phosphatase (0.5 μg/mL, Roche) in blocking solution overnight at 4°C. After washing in PBS with 0.1% Tween, sections were incubated in NBT/BCIP substrate solution at room temperature for 2–16 h for colorimetric detection. Protein concentration was determined using BCA assay (Thermo Scientific). For each sample, 10 μg of total protein were separated on SDS NuPAGE Bis-Tris 4–12% gels (Life Technology) in MES running buffer (Life Technology) and transferred to polyvinylidene fluoride membrane (Life Technology). The blots were blocked in 5% bovine serum albumin in PBS and 0.1% Tween 20 for 1 h at

**Western blot**

Cervical and thoracic DRGs were dissected on ice and quickly frozen in −80°C. Samples were removed into a microcentrifuge tube containing ice-cold sample buffer (20 mM Tris-HCl [pH 7.4], 1 mM dithiothreitol, 10 mM NaF, 2 mM Na3VO4, 1 mM EDTA, 1 mM EGTA, 5 mM microcin-LR, and 0.5 mM phenylmethylsulfonyl fluoride), and homogenized by sonication. Homogenates were centrifuged at 12,000 x g for 30 min at 4°C. The supernatant was used for analysis. Protein concentration was determined using BCA assay (Thermo Scientific). For each sample, 10 μg of total protein were separated on SDS NuPAGE Bis-Tris 4–12% gels (Life Technology) in MES running buffer (Life Technology) and transferred to polyvinylidene fluoride membrane (Life Technology). The blots were blocked in 5% bovine serum albumin in PBS and 0.1% Tween 20 for 1 h at
room temperature and incubated with rabbit anti-GRP (ImmunoStar, 1:5000), or rabbit anti-Actin (Sigma, 1:50,000) for 16 h at 4°C. This was followed by 1 h incubation in donkey horseradish peroxidase-linked secondary antibodies (Santa Cruz, 1:2500). Immunoblots were developed with the enhanced chemiluminescence reagents (Thermo Scientific). Band intensities were measured using Kodak 1D (version 3.6) and Actin served as internal control for normalization.

Reverse transcription polymerase chain reaction

RT-PCR was performed as previously described. Mice were sacrificed. DRG and spinal cord tissues were quickly dissected and rapidly frozen in dry ice. Total RNA was isolated and genomic DNA was removed in accordance with manufacturer’s instructions (RNeasy plus mini kit; QIAGEN). Single-stranded cDNA (total of 20 μL per sample) was synthesized from 1 μg RNA by using High Capacity cDNA Reverse Transcription Kit (Life Technologies). Gene expression of Grp and Tac1 was determined by real-time PCR (StepOnePlus; Applied Biosystems). Specific intron-spanning primers were designed with the NCBI Primer-BLAST. The primers used are:

- **Actb** (NM_007393.3): 5'-TGTTACCAACTGGGACGACA-3'; 5'-GGGGTGTTGAAGGTCTCAAA-3'; amplicon size: 166 bp.
- **Gapdh** (NM_008084.2): 5'-CCCAGCAAGGACACTAGCCAA-3'; 5'-TTATGGGGGTCTGGGATGGAAA-3'; amplicon size: 93 bp.
- **Grp** (NM_175012.3): 5'-TGGGCTGTGGGACACTTAAT-3'; 5'-GCTTCTAGGAGGTCCAGCAAA-3'; amplicon size: 146 bp.
- **Tac1** (NM_009311.2): 5'-GTGGCCCTGTTAAAGGCTAC-3'; 5'-TGCCCATTAGTCCAAACAGGA-3'; amplicon size: 85 bp.

Real-time PCR was carried out with FastStart Universal SYBR Green Master (Roche Applied Science). All samples were assayed in duplicates. PCR (heating at 95°C for 10 s and 60°C for 30 s) were performed. Data were analyzed using Comparative CT Method (StepOne Software v2.2.2.), and the expression of target mRNA was normalized to the expression of Actb and Gapdh.

RT-PCR with 2% agarose gel electrophoresis was performed using Taq DNA Polymerase (New England Biolabs) with 2 μL cDNA as a template per reaction. Reactions were optimized by running annealing temperature and cycle gradients with no-RT (∆RT) and no-template (H2O) samples as negative controls.

The following intron-spanning primer pairs and PCR parameters were used:

- **Actb**: 5'-GATGACGATATCGCTCGCTGGTCG-3'; 5'-GCCCTGTGGTACCAACAGGCCATACA-3'; amplicon size 447 bp. Parameters: 95°C 5 min, [95°C 40 s, 55°C 40 s, 72°C 40 s] for 21 cycles, 72°C 10 min.
- **Grp**: 5'-AGTCGAGAGCTCTGAGGGTT-3'; 5'-CCCTTGTCGTGTCTCCTCAAGCCTTTAACAGGGC-3'; amplicon size 329 bp. Parameters: 95°C 5 min, [95°C 40 s, 55°C 40 s, 72°C 40 s] for 28 cycles, 72°C 10 min.
- **Tac1**: 5'-GAGAGCAAAGAGCAGCGCCCCACG-3'; 5'-AAGAGCTTATAACAGGCCCAGG-3'; amplicon size 329 bp. Parameters: 95°C 5 min, [95°C 40 s, 55°C 40 s, 72°C 40 s] for 24 cycles, 72°C 10 min.

RNA-seq library preparation

Three adult male C57BL/6J mice were sacrificed by decapitation. Cervical and thoracic DRGs were dissected out and snap frozen on dry ice. Total RNA was extracted using RNeasy Plus Micro Kit (Qiagen) following manufacturer’s instructions. RNA concentration and quality was examined using Agilent Bioanalyzer Chip. RNA integrity numbers were above 8.50. Library preparation was performed with Epicentre Ribozero Gold kit (Clontech) according to manufacturer’s protocol. cDNA was then blunt ended, an A base was added to the 3’ ends, followed by the ligation of Illumina sequencing adapters to the ends. Ligated fragments were then amplified for 12 cycles using primers incorporating unique index tags. Fragments were sequenced on an Illumina HiSeq-2500 using single reads extending 50 bases. Sequencing depth was 55-60M reads per sample.

RNA-seq data acquisition, quality control, and processing

RNA-seq reads were aligned to the GRCm38.76 assembly from Ensembl with STAR version 2.0.4b. Gene counts were derived from the number of uniquely aligned unambiguous reads by Subread:featureCount version 1.4.5. Transcript counts were produced by Sailfish version 0.6.3. Sequencing performance was assessed for total number of aligned reads, total number of uniquely aligned reads, genes and transcripts detected, ribosomal fraction known junction saturation and read distribution over known gene models with RSeQC version 2.3. All gene-level and transcript counts were then imported into the R/Bioconductor package EdgeR and TMM normalized to adjust for differences in library size. Genes or
transcripts not expressed in any sample were excluded from further analysis. Performance of the samples was assessed with a spearman correlation matrix and multi-dimensional scaling plots. Generalized linear models with robust dispersion estimates were created to test for gene/transcript level differential expression. The fit of the trended and tagwise dispersion estimates were then plotted to confirm proper fit of the observed mean to variance relationship where the tagwise dispersions are equivalent to the biological coefficients of variation of each gene. Differentially expressed genes and transcripts were then filtered for FDR adjusted p values less than or equal to 0.05.

To enhance the biological interpretation of the large set of transcripts, grouping of genes/transcripts based on functional similarity was achieved using the R/Bioconductor packages GAGE and Pathview. GAGE and Pathview were also used to perform pathway maps on known signaling and metabolism pathways curated by KEGG.

**GRP protein sequences**

Amino acid sequences were obtained from NCBI nucleotide database. The following Genbank Accession numbers are for each species:

- **Mouse**: NM_175012.4
- **Rat**: NM_133570.5
- **Chimpanzee**: XM_001142106.3
- **Human**: NM_002091.3
- **Dog**: XM_861026.3
- **Rabbit**: XM_003474134.3
- **Guinea Pig**: NM_001090321.1
- **Cow**: NM_001101239.1
- **Sheep**: XM_00279900.1
- **Horse**: XM_001489303.1
- **Chicken**: XM_001277900.1

**Statistics**

Values are reported as the mean ± standard error of the mean (SEM). Statistical analyses were performed using Prism 6 (v6.0e, GraphPad, San Diego, CA). For comparison between two groups, unpaired or paired two-tailed t test was used. Normality and equal variance tests were performed for all statistical analyses. p < 0.05 was considered statistically significant.

**Results**

**GRP antibody specifically detects GRP in DRGs and the spinal cord**

GRP is initially translated as a 148 amino acid prepropeptide that is further processed into smaller biologically active peptides comprising 27 (GRP1-27) or 10 amino acids (GRP18-27). Comparison of GRP18-27 across species reveals that GRP is highly conserved across many species with no differences in the last seven amino acids (WAVGHLM) (Table 2), suggesting that this region is important for its function.

### Table 2. Comparison of amino acid sequences of GRP18-27 across species.

| Species   | Amino Acid Sequence       |
|-----------|---------------------------|
| Mouse     | GSHWAVGHLM-amide          |
| Rat       | GSHWAVGHLM-amide          |
| Chimpanzee| GNHWAVGHLM-amide          |
| Human     | GNHWAVGHLM-amide          |
| Guinea Pig| GNHWAVGHLM-amide          |
| Cow       | GNHWAVGHLM-amide          |
| Sheep     | GNHWAVGHLM-amide          |
| Horse     | GNHWAVGHLM-amide          |
| Chicken   | GSHWAVGHLM-amide          |

Blue amino acids are conserved across different peptides or across species. Red mino acids indicate differences.

### Table 3. Comparison of amino acid sequences of mouse neuropeptides.

| Peptide   | Amino Acid Sequence       |
|-----------|---------------------------|
| GRP18-27  | GSHWAVGHLM-amide          |
| Bombesin  | pEQRLGNQWAVGHLM-amide     |
| NeuromedinB| GNLUWATGFM-amide        |
| Substance P| RPKPQQFFGLM-amide       |

Green amino acids are conserved across different peptides. Red amino acids indicate differences.

Comparison of GRP18-27 with bombesin (toad), as well as neuromedin B (NMB) and SP, reveals some distinct regions (Table 3). The majority of the GRP antibodies are raised against the seven amino acids WAVGHLM which are identical between GRP and bombesin. A recent study suggested that the GRP antibody is only specific at high dilution (1:4000), and GRP could not be detected in DRGs. To address the discrepancies between this study and the others, we first repeated IHC using the GRP antibody at 1:500 and 1:1000 dilutions. Consistent with our previous studies, at both dilutions, GRP was clearly detected in wild type (WT) dorsal horn (Figure 1(a) and (c)) but nearly absent in Grp knockout (KO) (Figure 1(b) and(d)). However, at 1:4000, GRP was nearly undetectable in WT dorsal horn compared to lower dilutions (Figure 1(e)), while staining was again absent in Grp KO (Figure 1(f)). In DRG tissues, results were similar to dorsal horn. At 1:500 (Figure 1(g) and (h)) and 1:1000 (Figure 1(i) and (j)), GRP+ DRG cell bodies were clearly visible in WT but absent in Grp KO. At 1:4000 (Figure 1(k) and (l)), GRP+ DRG cell bodies were not detectable in either WT or Grp KO. Based on these results, we conclude that the GRP antibody is specific and capable of detecting GRP in DRG and spinal cord tissues. Our results from different antibody dilutions, though, may vary with different production lots. For
all further staining experiments in this study, we used the 1:1000 dilution.

Many studies have examined GRP expression by IHC, whereas Western blot has not been used. Next, we examined the specificity of the GRP antibody using Western blot analysis of DRG protein homogenates (Figure 1(m)). Although this antibody recognizes several non-specific large sized bands, an approximately 16 kDa band that is equivalent to the size of the full length of GRP prepro-protein was detected in WT but absent in Grp KO (Figure 1(m), red arrow). In Grp KO tissues, a faint band was visible, but its molecular weight is slightly larger than GRP band in WT and thus considered to be non-specific. These results indicate that the GRP antibody from Immunostar can be used to specifically detect GRP in DRGs by Western blot.

Ablation of primary afferent terminals in the dorsal horn eliminates most of GRP protein

Conflicting results have been reported concerning the expression of GRP protein in DRGs: Fleming et al.28 showed that the GRP antibody is specific at 1:500 dilution with approximately 2% of DRGs positive for GRP, while Solorzano et al.25 indicated otherwise. Moreover, an earlier study showed that intrathecal capsaicin injection not only ablated TRPV1+ afferents but also ablated GRP+ afferents in the dorsal horn,5 whereas the Solorzano et al.25 study did not indicate any difference in GRP staining in the dorsal horn following capsaicin injection.

To further investigate the origin of GRP in dorsal horn, mice were intrathecally injected with

Figure 1. Specific detection of GRP in spinal cord dorsal horn and DRG tissues. (a)–(f) Lumbar dorsal horn IHC images of GRP antibody staining at different dilutions. At 1:500 dilution, GRP is clearly detected in WT (a) but nearly absent in Grp KO (b) littermates. At 1:1000 dilution GRP is still detected in WT (c) but absent in Grp KO (d). However, at 1:4000 dilution GRP is barely detected in WT (e) and still absent in Grp KO (f). (g)–(l), DRG IHC images of GRP antibody at varying dilutions. At 1:500 dilution, GRP+ DRG neuron cell bodies are clearly detected in WT (g) but almost absent in Grp KO (h) littermates. At 1:1000 dilution GRP+ neurons are still detected in WT (i) but absent in Grp KO (j). However, at 1:4000 dilution GRP+ neurons are barely detected in WT (k) and still absent in Grp KO (l). (m) Western blot showed that GRP antibody (1:5000) recognized a band at approximately 16 kDa in WT DRGs, which was not detected in Grp KO DRGs. A nonspecific band (~30 kDa) was also shown in all samples. (a)–(l) n = 3 mice per genotype and 10 sections per group. (m) n = 2 mice per genotype and 20–24 DRGs per animal. Scale bar = 100 μM.
resiniferatoxin (RTX), a potent TRPV1 agonist, to ablate TRPV1+ afferents. Previous studies have demonstrated that RTX reliably ablates TRPV1+ afferents, while not affecting TRPV1+ DRG cell bodies. Because approximately 80% of GRP+ DRG neurons express TRPV1, RTX-injection should abolish most of the GRP staining in dorsal horn. Quantitative analysis of TRPV1 staining confirmed almost complete ablation of TRPV1+ afferents in dorsal horn of RTX-injected mice compared to saline-injected control (Figure 2(a)–(c)). Similarly, GRP staining within the same section was nearly abolished in the dorsal horn of RTX-injected mice (Figure 2(d)–(f)). Double IHC images showed an overlap in expression of TRPV1 and GRP in control mice that is eliminated by RTX ablation of TRPV1+ afferents (Figure 2(g) and (h)).

Four independent groups have recently used dorsal rhizotomy of the lumbar spinal cord to evaluate the origin of GRPergic fibers in mice and rats and reported conflicting results.13,16,18,25,28 Given the inconsistencies and the importance of a complete surgery, it is important to show images of the entire section of the lumbar dorsal horn, both ipsilateral and contralateral sides, a gold standard in the field. It is worth noting that two of these studies that supported the majority of GRP as being derived from the dorsal horn did not show the entire section of the spinal cord, making their results difficult to evaluate and interpret.25,28 In contrast, we and other researchers showed consistent findings in the spinal cord of mice and rats.13,18 To further improve the quality of immunostaining for evaluation and to see if our previous finding could be reproduced, one person from our lab performed unilateral rhizotomy, whereas another person carried out double IHC studies on the spinal cord sections. CGRP staining demonstrated almost complete ablation of CGRP+ afferents in the ipsilateral dorsal horn compared to contralateral (Figure 2(i) and (j)), suggesting efficient rhizotomy. Consistent with recent studies in mice and rats,13,18 GRP staining within the same section was also nearly abolished (Figure 2(k) and (l)). Merged images (Figure 2(m)). Taken together, the RTX-mediated TRPV1 fiber ablation and rhizotomy results support the notion that most of GRP, as well as TRPV1 and CGRP, in the dorsal horn is of primary afferent origin. It cannot be excluded, though, that small amounts of these peptides and channels in the dorsal horn may originate from spinal cord interneurons or from other sources such as descending projections. For example, previous studies suggested that GRP is expressed in neurons of the lateral parabrachial nucleus, a region which has been shown to project directly to the spinal cord dorsal horn.

**GRP is detectable in DRG, but not dorsal horn, primary cultures**

To examine if there are GRP+ neurons in the dorsal horn, we dissected out DRGs and dorsal horn of the spinal cord, acutely dissociated DRG and dorsal horn neurons and cultured the neurons for GRP and NeuN double IHC (Figure 3(a)). We found that about 3.8% (6/157) of NeuN+ DRG neurons were GRP+ (Figure 3(b)). However, we were unable to find any GRP+ dorsal horn neurons (Figure 3(b)). These results suggest that GRP protein is not expressed in dorsal horn neurons or the expression level falls below the detectable limit.

**GRP expression partially, but not completely, overlaps with substance P in DRG neurons**

Although GRP and NMB share five of the seven amino acids in their C-terminal regions (Table 3), several pieces of evidence indicate that the GRP antibody does not recognize NMB: despite overlapping expression, NMB is also detected in non-peptidergic neurons.40 In contrast, SP only has homology with GRP in the last two amino acids (Table 3), with the last one, methionine (M) shared by many neuropeptides. Thus, it is unlikely that one amino acid would make the GRP antibody recognize only SP even though there is cross-reactivity. Indeed, it has been demonstrated that GRP and SP antibodies also recognized neurons singularly expressing either SP or GRP.

We revisited this issue by performing double IHC on DRG neurons from WT and Grp KO mice (Figure 4). Both GRP (Figure 4(a)) and SP (Figure 4(c)) were detected in the subset of WT DRG neurons. In contrast, while GRP was absent in Grp KO mice (Figure 4(b)), SP expression was not affected in Grp KO mice (Figure 4(d)). Merged images (Figure 4(e) and (f)) show that about 50% of GRP+ neurons are SP+ in WT DRG. A Venn diagram further illustrates the GRP and SP neuron overlap and percentages in WT DRGs (Figure 4(g)). Consistent with previous studies,13,16 the percentage of GRP+ neurons was approximately 8%, whereas the percentage of SP+ neurons was approximately 12.5% of the total.

**GRP+ and SP+ terminals make contacts with GRPR+ neurons in dorsal horn**

To examine whether SP+ or GRP+ terminals make connections with GRPR+ neurons in the spinal dorsal cord, triple IHC was performed for SP, eGFP, and GRP in GRPR-eGFP mice. Some GRPR+ neurons and densely SP+ and GRP+ terminals were found in the superficial part (laminae I and IIo) of the spinal dorsal horn (Figure 5(a)–(c)). Many SP+ or GRP+ fibers could be observed making contacts with GRPR+ neurons.
Figure 2. Chemical or surgical ablation of primary afferent terminals eliminates most of GRP protein in the dorsal horn. (a)–(c) Lumbar dorsal horn IHC images of TRPV1 staining from mice injected intrathecally with saline (a) or RTX (25 ng) (b) and normalized intensities (c) indicate near complete ablation of TRPV1 signal in RTX-treated mice. (d)–(f) Comparison of GRP staining images from saline (d) and RTX (e) groups and intensities (f) also show almost total ablation of GRP signal in RTX-treated mice. (g, h) Merged images indicate most GRP staining overlaps with TRPV1 in saline control (g) which is mostly absent in RTX-treated mice (h). (i, j) A single lumbar dorsal horn image of CGRP staining from dorsal rhizotomy mice of both contralateral and ipsilateral sides of the surgery (i) and normalized intensities of each side (j) shows that rhizotomy eliminated nearly all CGRP signal in the ipsilateral side. (k, l) GRP staining image of rhizotomy contra- and ipsilateral dorsal horn (k) and intensities (l) indicates an almost complete ablation of GRP signal in ipsilateral side compared with contralateral. (m) Merged image shows significant overlap in GRP and CGRP staining in contralateral side that is eliminated in ipsilateral side following rhizotomy. Scale bar = 100 μM. Data are presented as mean ± SEM. n = 3 mice per group and 10 sections per group. Unpaired t test in (c) and (f). Paired t test in (j), and (l), ***p < 0.001.
(Figure 5(d)–(f)), and a few SP and GRP double-labeled terminals could be found making contacts with GRPR<sup>+</sup> neurons (Figure 5(f)).

To examine the colocalization of SP and GRP in the spinal dorsal horn, double immuno-electron microscopic (Immuno-EM) for SP and GRP were performed in the lumbar cord. Some SP<sup>+</sup> terminals identified by the silver-enhanced nanogold particles enhancement are colocalized with GRP<sup>+</sup> terminals revealed by the immunoperoxidase reaction products (Figure 5(g)), and some axon terminals only express GRP (Figure 5(h)) or SP (not shown). SP<sup>+</sup> or GRP<sup>+</sup> terminals formed asymmetric synapses with GRPR<sup>+</sup> dendritic profiles (Figure 5(i)–(k)), and more GRP<sup>+</sup> fibers formed synaptic connections with GRPR<sup>+</sup> neurons than SP<sup>+</sup> fibers (~2:1 GRP:SP contacts with GRPR neurons). Although a majority of GRP<sup>+</sup> fibers in the dorsal horn are of primary afferent origin, it cannot be excluded that some of the GRP-GRPR and SP-GRPR contacts in the dorsal horn may be due to GRP<sup>+</sup> or SP<sup>+</sup> descending projections from the brain as mentioned earlier.

Genetic deletion of Tac1 results in reduced peptidergic expression in spinal cord dorsal horn

The Tac1 gene encodes the propeptide that is cleaved to generate SP and three other neuropeptides of the tachykinin family. Solorzano et al. showed that GRP expression is reduced in the spinal cord of Tac1 KO mice, which was interpreted as evidence for a presumed lack of specificity of the GRP antibody. Because Tac1 KO mice also exhibited deficits in expression of other antibody immunostaining, we speculated that an attenuated GRP expression may reflect broader abnormalities of gene expression in the primary afferents of Tac1 KO mice.
To examine whether lack of SP may influence expression of other molecular markers, we performed IHC staining in spinal dorsal horn sections from WT and Tac1 KO littermates. As expected, SP staining was absent in Tac1 KO dorsal horn (Figure 6(a)–(c)). Consistent with previous results, GRP staining was significantly reduced in Tac1 KO compared to WT (Figure 6(d)–(f)). Remarkably, a small reduction, yet significant, was observed for CGRP staining (Figure 6(g)–(i)), and NPY staining was dramatically reduced (Figure 6(j)–(l)) in Tac1 KO relative to the control. However, the non-peptidergic IB4-binding pattern and intensity was similar in WT and Tac1 KO dorsal horn (Figure 6(m)–(o)). Taken together, these results suggest that Tac1 is important for either expression and/or trafficking of neuropeptides in peptidergic primary afferents.

**Optimized methods for detection of GRP in DRG**

Considerable variations of GRP immunostaining have been observed in DRG neurons, ranging from somewhat widespread GRP staining to more distinct or restricted staining. One study even described two types of GRP neurons comprising low and high level of expression. We compared two different staining methods: either mounted on slides prior to antibody incubation, or sections stained floating freely in PBS, and mounted after staining. Using the same antibody, on-slide staining resulted in a widespread staining pattern for GRP (Figure 7(a)), whereas free-floating staining of DRG sections revealed a more distinct and specific pattern (Figure 7(b)). These results suggest that the discrepancies can be attributed to differences in IHC protocols rather than insufficient antibody specificity.

Concerning Grp mRNA expression in DRG, a few studies reported negative ISH results, while our recent study showed detectable ISH signal for Grp in WT DRG neurons, which is comparable to GRP IHC staining pattern. To optimize the methods for reliable Grp mRNA detection, we tested different incubation times for color development of Grp ISH signals on WT DRG sections. Grp signals were barely detectable after 4 h of incubation in NBT/BCIP colorimetric substrate (Figure 7(c), arrows), but continuing incubation to 16 h produced specific Grp signals (Figure 7(d), arrowheads). The fact that detection of Grp signals by ISH requires a longer incubation time for color development than most other probes suggests that Grp transcripts are likely present in low copy number, rendering them difficult to detect by conventional protocols.

We next performed molecular expression analyses by RNA-seq in DRGs from adult mice. Read length was set at 50 bases with a sequencing depth in a range of 55–60 million single end reads to allow detection of low-abundance transcripts such as Grp, according to the ENCODE Consortium’s “Standards, Guidelines and Best Practices for RNA-seq” (https://genome.ucsc.edu/ENCODE/protocols/dataStandards/ENCODE_RNAseq_Standards_V1.0.pdf). Transcript abundance is expressed as FPKM (Fragments Per Kilobase of transcript per Million fragments mapped), which provides a length and depth normalization to permit comparisons both within and between samples. In this condition, we detected the level of Grp transcripts in WT DRGs as 0.27 ± 0.0033 FPKM, which is lower than the level of Npy (1.91 ± 0.47 FPKM), Mrgpra3 (3.87 ± 0.85 FPKM), Tac1 (18.45 ± 2.42 FPKM), and Nmb (48.81 ± 9.33 FPKM) but very close to the level of Pdyn.
GRP transcripts in mouse DRGs can be detected by RNA-seq. Moreover, its expression is even lower than Npy, a gene whose mRNA expression often falls below the threshold of detection by conventional ISH technique.

Finally, we performed RT-PCR followed by gel electrophoresis to further analyze the abundance of Grp transcripts in intact DRGs, as Grp became detectable only at \( \geq 32 \) cycles (Figure 7(e)), whereas Nppb, which encodes B-type neuropeptide and Actb were observed after 26 and 18 cycles, respectively (Figure 7(e)).
finding is consistent with previous studies\textsuperscript{28,30} and confirmed the presence of Grp mRNA in uncultured DRGs. It also underscores the fact that sufficient PCR cycles are required for Grp detection.

**Increased numbers of Grp\textsuperscript{+}, but not Tac1\textsuperscript{+}, neurons and upregulation of Grp mRNA levels in dry skin-induced chronic itch DRGs**

SP-NK1R signaling has been implicated in itch transmission and development of chronic itch.\textsuperscript{23,50,51} However, it remains unclear whether Tac1/SP expression in DRG increases in chronic itch conditions. Therefore, we used the AEW dry skin mouse model of chronic itch model to assess the effects of chronic itch on Tac1 mRNA expression in DRG. Grp expression was absent in Grp KO neurons (Figure 8(a)) demonstrating that the probe is specific for Grp. Consistent with our previous studies,\textsuperscript{12} the percentage of Grp\textsuperscript{+} neurons was almost doubled in DRG from dry skin mice compared to water-treated control (Figure 8(b)–(d)). In contrast, the percentage of Tac1\textsuperscript{+} neurons, which was unaffected in Grp KO (Figure 8(e)), was comparable in DRG between dry skin and the control mice (Figure 8(f)–(h)). Lastly, we performed RT-PCR with gel electrophoresis to visualize Grp and Tac1 mRNA expression with Actb as control, which clearly showed increased band intensities for Grp in dry skin mice, whereas Tac1 in dry skin appeared similar to control (Figure 8(i)). qRT-PCR was also performed that showed Grp

**Figure 6.** Apparent reduction of peptidergic markers in dorsal horn of Tac1 gene-deleted mice. (a)–(c) Lumbar dorsal horn IHC images of SP staining in WT (a) and Tac1 KO (b) and normalized intensities (c) show loss of SP expression. (d)–(f) IHC images of GRP staining in WT (d) and Tac1 KO (e) with normalized intensities (f) indicate a significant reduction in GRP signal in Tac1 KO. (g)–(i) CGRP staining images in WT (g) and Tac1 KO (h) with intensities (i) also show a small, but significant, reduction in CGRP in Tac1 KO. (j)–(l) NPY staining images in WT (g) and Tac1 KO (h) with intensities (i) indicates a large reduction in NPY signal in Tac1 KO. (m)–(o) IB4-binding images in WT (m) and Tac1 KO (n) with intensities (o) show no apparent differences. Scale bar = 100 μM. Data are presented as mean ± SEM. n = 3 mice per genotype and 10 sections per group. Unpaired t-test in (c), (f), (i), and (l), *p < 0.05, **p < 0.01, ***p < 0.001.
Dry skin-induced chronic itch in mice increases the numbers of GRP, but not SP, positive neurons in the DRG

Next, double IHC was performed to examine SP and GRP expression in DRG sections of mice with dry skin.
itch. Compared to water-treated control (Figure 9(a)), the percentage of GRP\(^+\) neurons in DRG was nearly doubled in dry skin-induced chronic itch (Figure 9(b) and (c)), consistent with the results from Grp\(\)ISH.

Also consistent with Tac1\(\)ISH, the percentage of SP\(^+\) neurons was similar in dry skin-induced chronic itch DRG compared to control (Figure 9(d)–(f)). Merged images (Figure 9(g) and (h)) indicate no significant difference in the percentage of double-positive neurons in dry skin mice compared to control (Figure 9(i)) suggesting ectopic expression of GRP in non-SP DRG neurons in chronic itch. To further confirm the immunostaining results, Western blot of DRG protein extracts was performed using the GRP antibody (Figure 9(j) and (k)). Western blot of dry skin DRG protein samples indicated an apparent upregulation of GRP levels compared to control (Figure 9(j)), and quantitative analysis showed significant increase of GRP in dry skin DRGs (Figure 9(k)). Taken together, these data confirm previous results that GRP protein is upregulated in chronic itch,\(^{12,13}\) whereas SP is not increased in chronic itch. There are conflicting reports of SP expression related to chronic itch. Some studies reported increases in SP skin fibers and mast cells in human atopic dermatitis (AD) skin and mouse AD models,\(^{51-53}\) whereas other studies found reduced SP in AD skin.\(^{54,55}\) However, to our knowledge, no other studies have investigated SP expression in DRG in dry skin-induced chronic itch conditions. Our findings suggest that in dry skin-induced chronic itch, SP expression in DRG is largely unaffected.

**Neither Grp nor Tac1 expression are increased in dorsal horn of mice with dry skin-induced chronic itch**

While it is clear that Grp expression in DRG is upregulated during chronic itch, the effect of chronic itch on Grp expression in the dorsal horn has not been investigated. This issue is important because, assuming Grp\(^+\) dorsal horn neurons are important for chronic itch, one would anticipate an upregulation of Grp mRNA in the spinal cord of mice with dry skin itch. To examine this, we performed Grp and Tac1\(\)ISH and found that the
number of Grp$^+$ neurons in both cervical and thoracic dorsal horn was similar in dry skin mice and controls (Figure 10(a)–(f)). Tac1$^+$ neuron numbers also appeared unchanged in both cervical and thoracic dorsal horn from dry skin mice compared to controls (Figure 10(g)–(l)). qRT-PCR of cervico-thoracic spinal cord cDNA also revealed no significant differences in Grp (Figure 10(m)) or Tac1 (Figure 10(n)) expression in dry skin mice compared to control.

**Discussion**

The specificity of the GRP antibody and cross-reactivity with SP

In this study, we show that different IHC protocols could have major effects on the staining outcome even though the same GRP antibody was used. Regardless of the method used, optimization of the procedure is a pre-requisite for performing specific GRP immunostaining,
as many pitfalls such as perfusion and the quality of tissues may influence the outcome of IHC. Although the GRP antibody is specific, it remains one of the most difficult antibodies to work with in our experience. This is also true for Grpr ISH, as many investigators are unable to detect Grpr in the spinal cord. Thus, if distinct GRP immunostaining cannot be achieved in DRG, it would be difficult to interpret the results obtained from spinal cord immunostaining or double IHC staining (GRP vs. SP).

Our double IHC using GRP and SP antibodies is consistent with studies in rats, as well as supported by EM analysis revealing contacts not only between SP⁺ fibers but also GRP/SP fibers and GRPR neurons. Although SPergic fibers form contacts with GRPR neurons, it is unlikely for SP to communicate with GRPR neurons because they appear to lack NK1 receptor. Interestingly, capsaicin treatment could induce SP, but not GRP, release from spinal cord slices. Conversely, it is conceivable that GRP is selectively released from SP/
GRPergic fibers, but SP does not, in response to pruritogenic stimuli. It is also possible that SP in GRPergic fibers is selectively used to relay itch via NK1R neurons.23

**Grp mRNA and GRP protein expression in the spinal cord**

Of four recent independent investigations of the origin of GRP in the spinal cord using the dorsal rhizotomy, two studies, displaying both ipsilateral and contralateral side of the spinal cord in their entirety in one image, produced highly consistent results.15,18 The other two, which argue for endogenous expression of GRP in the spinal cord, in fact were contradictory in respect to their GRP immunostaining in DRGs.25,28 As aforementioned, a reasonable explanation for inconsistencies is likely due to the experimental protocols used rather than antibody specificity. More recently, we found that specific GRP immunostaining could be obtained using fresh DRG tissue without perfusion (data not shown). Notably, the finding that detection of Grp mRNA is refractory to ISH is reminiscent of the absence of Npy mRNA signal as examined by ISH in DRG.58

It has been known that not all mRNAs are translated into protein and a gene transcript could be translated into protein in one tissue but not the other. For example, although temporal and spatial expression pattern of Grp mRNA strikingly resembles that of GRP protein in lung tissue, some Grp+ tissues were negative for GRP immunostaining.59–63 Moreover, our unpublished data also indicate that in the brain, not all Grp mRNA-expressing areas are positive for GRP immunostaining. However, we are unable to exclude with certainty that GRP, upon translated by dorsal horn neurons, is rapidly degraded or GRP intrinsic to the dorsal horn falls below the threshold of IHC method we used. Such a possibility, however, is still compatible with the conclusion that a majority of GRPergic fibers in the spinal cord are of peripheral origin. Moreover, the present study further supports the notion the remaining Grp+ fibers in the dorsal horn after the dorsal rhizotomy are perhaps of descending origin, because of the absence of Grp+ immunostaining in dissociated dorsal horn neuronal culture. Several studies have used the Grp-eGFP line which lack eGFP in DRGs as one of the evidence to argue that GRP protein is abundantly expressed in the dorsal horn but not in DRGs.25,30,64 As we noted previously,12 eGFP could be expressed even in knock-out mice with an eGFP knock-in. Conversely, eGFP in transgenic or knock-in mice may not be expressed in the regions where endogenous mRNA or protein of the gene of interest is present. While Grp-eGFP mice largely recapitulate endogenous Grp mRNA expression in the spinal cord, it is worth noting that expression of eGFP reporter protein only indicates eGFP mRNA transcription (thereby a surrogate for Grp mRNA), but the translation of the eGFP protein from eGFP mRNA cannot be used for an indication of the translation of GFP protein from Grp mRNA, because endogenous Grp mRNA, versus the eGFP mRNA, may be subject to distinct translational control.

**Detection of Grp mRNA by RT-PCR, ISH, and RNA-seq**

The present study indicates that the copy number of the Grp transcript is much lower than Nmb, Tac1, and Mrgpra3 (Table 4). Thus, sufficient PCR cycles and incubation times for color development by ISH are also crucial for signal detection. Although both Grp and Mrgpra3 are expressed in small percentage of DRGs, the level of Grp expression is approximately 90% lower than that of Mrgpra3 according to RNA-Seq. This explains why it is easier to detect Mrgpra3 than Grp. Although RNA-seq has been widely used for high-throughput profiling of gene expression, the technique could be limited by its inability to detect rare or low abundance of gene transcripts.65,66 A key challenge is to increase the cell number rather than deeper sequencing.67 For example, a single-cell RNA-seq has recently been applied to DRG neurons, revealing remarkable detailed transcriptome in a single-cell resolution. The study, however, failed to detect Grp mRNA based on average of 50 individual cells.62 Given the low copy number of Grp transcripts expressed in very small percentage of DRGs, the reads for Grp mRNA could be zero or fall into the baseline noise if not enough cells are sampled. Other system errors which are inherently associated with RNA-seq may also skew the value for Grp mRNA.68 To increase mRNA capture efficiency and overcome the limitation of single cells that can be sequenced, a droplet platform has been recently developed, enabling deep sequencing of a large number of cells instead of dozens or hundreds at a time.69 In this regard, one may use Grp as a positive control to evaluate the sensitivity and coverage of the gene expression profiling with rare and low-copy number by RNA-seq.

**The role of Grp mRNA vs. protein**

The debate on the expression of GRP raises several interesting issues that are worth considering here. Concerning the specificity of the GRP antibody, a review of recent literature indicates that several groups were able to obtain specific GRP immunostaining in DRGs and their results are highly consistent (Table 1). Despite the demonstrated specificity of the GRP antibody, detection of distinct GRP immunostaining in subsets of DRGs remains challenging, as optimization of the immunostaining condition could be time-consuming. Some
Barry et al.

researchers may be unwilling to spend weeks or even months on this. By contrast, it would be much easier to obtain widespread immunostaining using the GRP antibody if the condition is not optimized. Based on our own experience, detection of Grp positive signaling in DRGs by ISH is even more daunting, simply because of the low copies of Grp mRNA. This raises interesting questions that have puzzled some investigators. First, why is Grp mRNA expressed in the dorsal spinal cord if it is physiologically unimportant? One possibility is that spinal Grp may reflect an ontogenetic/evolutionary relic. Since the neural crest that gives rise to DRGs emerges from the dorsal neural tube, many peptide genes in DRGs might have co-opted from the dorsal spinal cord and diverged to adopt novel functions, whereas their roles in the spinal cord became diminished. This enables the rapid and simple transduction of modality-specific information from the periphery to the spinal cord. Similarly, many postsynaptic receptors in the spinal cord crucial for relaying information encoded by primary afferents-released peptides may still reside in DRG, but they may have minimal functionality. Second, why is Grp/Grpr mRNA expression so low relative to some other genes if they are important itch genes? In DRGs, existing data do not suggest a positive correlation between abundance of a peptide and its physiological importance. For example, Nppb is more abundantly expressed than Grp in DRGs, but at least 10 times higher amount of BNP is required to evoke scratching behavior similar to that by GRP, even though the time course of the slow onset of scratching behavior does not support a direct activation of NPRA by BNP in the spinal cord. From an evolutionary perspective, GRP may have been adopted to alert the body of pruritogenic stimuli. Given the relatively high affinity of GRP binding to its receptor, a transient and rapid release of GRP protein stored at the terminals of primary afferents should be sufficient to activate GRPR to evoke a few bouts of scratches, sub serving the warning mechanism. By contrast, a persistent and large amount of GRP release may trigger vicious scratch-itch cycles, resulting in a pathological itch condition. As such, there is a need to minimize Grp mRNA in DRGs until GRP at the terminal is largely depleted. Similar to Grp, NPY peptide is present in the nerve terminals, but the level of its mRNA expression is very low in DRGs according to RNA-Seq, limiting its detection by ISH. Thus, the level of Grp mRNA cannot be equated to that of GRP protein. One can envision that upon acute pruritogenic stimuli, GRP protein at the terminals could be rapidly released, while Grp mRNA translation may proceed slowly to maintain the level of GRP protein synthesis at a normal physiological state.

In summary, using a combination of molecular, anatomic, genetic, ISH, IHC, Western blot, RNA-seq, and ultrastructure analysis approaches, we demonstrate the presence of GRP in primary afferents as well as lack of evidence for GRP protein synthesis intrinsic to dorsal horn neurons. The present study indicates that the disagreement on GRP expression in DRGs and the antibody specificity is likely due to different IHC protocols used as well as low abundance of Grp mRNA in DRGs.

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Author contributions

ZFC conceived the project and DMB, HL, YQL, and ZFC designed the experiments; DMB, HL, XYL, KFS, XTL, XJC, and JY performed experiments and data analysis. LH, ZYW, and YQL performed EM analysis; XJC and YGS conducted IHC independently, JY and AM contributed to the work and DMB, HL, XYL, YQL, and ZFC wrote the manuscript. DMB and HL contributed equally to this work.

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