Temporal sequence of activation of cells involved in purinergic neurotransmission in the colon

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Key points

- Platelet derived growth factor receptor α (PDGFRα+) cells in colonic muscles are innervated by enteric inhibitory motor neurons.
- PDGFRα+ cells generate Ca²⁺ transients in response to exogenous purines and these responses were blocked by MRS-2500.
- Stimulation of enteric neurons, with cholinergic and nitrergic components blocked, evoked Ca²⁺ transients in PDGFRα+ and smooth muscle cells (SMCs).
- Responses to nerve stimulation were abolished by MRS-2500 and not observed in muscles with genetic deactivation of P2Y1 receptors.
- Ca²⁺ transients evoked by nerve stimulation in PDGFRα+ cells showed the same temporal characteristics as electrophysiological responses.
- PDGFRα+ cells express gap junction genes, and drugs that inhibit gap junctions blocked neural responses in SMCs, but not in nerve processes or PDGFRα+ cells.
- PDGFRα+ cells are directly innervated by inhibitory motor neurons and purinergic responses are conducted to SMCs via gap junctions.

Abstract

Interstitial cells, known as platelet derived growth factor receptor α (PDGFRα+) cells, are closely associated with varicosities of enteric motor neurons and suggested to mediate purinergic hyperpolarization responses in smooth muscles of the gastrointestinal tract (GI), but this concept has not been demonstrated directly in intact muscles. We used confocal microscopy to monitor Ca²⁺ transients in neurons and post-junctional cells of the murine colon evoked by exogenous purines or electrical field stimulation (EFS) of enteric neurons. EFS (1–20 Hz) caused Ca²⁺ transients in enteric motor nerve processes and then in PDGFRα+ cells shortly after the onset of stimulation (latency from EFS was 280 ms at 10 Hz). Responses in smooth muscle cells (SMCs) were typically a small decrease in Ca²⁺ fluorescence just after the initiation of Ca²⁺ transients in PDGFRα+ cells. Upon cessation of EFS, several fast Ca²⁺ transients were noted in SMCs (rebound excitation). Strong correlation was noted in the temporal characteristics of Ca²⁺ transients evoked in PDGFRα+ cells by EFS and inhibitory junction potentials (IJP)s recorded with intracellular microelectrodes. Ca²⁺ transients and IJP s elicited by EFS were blocked by MRS-2500, a P2Y1 antagonist, and absent in P2ry1(−/−) mice. PDGFRα+ cells expressed gap junction genes, and gap junction uncouplers, 18β-glycyrrhetinic acid (18β-GA) and octanol blocked Ca²⁺ transients in SMCs but not in neurons or PDGFRα+ cells. IJP s recorded from SMCs were also blocked. These findings demonstrate direct innervation of PDGFRα+ cells by motor neurons. PDGFRα+ cells are primary targets for purinergic neurotransmitter(s) in enteric inhibitory neurotransmission. Hyperpolarization responses are conducted to SMCs via gap junctions.
Introduction

Gastrointestinal (GI) motility is regulated by the enteric nervous system, which includes excitatory and inhibitory motor neurons innervating the muscle layers (Burnstock et al. 1963; Bennett, 1966; Waterman & Costa, 1994; Spencer & Smith, 2001). Inhibitory neurotransmission reduces the excitability of smooth muscle cells (SMCs), reducing Ca^{2+} entry through voltage-dependent Ca^{2+} channels and relaxing muscles. Inhibitory motor neurons release nitric oxide (NO), purines, vasoactive intestinal polypeptide (VIP) and pituitary adenylate cyclase-activating polypeptide (PACAP) (Burnstock et al. 1980; Bult et al. 1990; Crist et al. 1992; Grider et al. 1994; Mutafova-Yambolieva et al. 2007). The post-junctional electrical response to enteric inhibitory neurotransmission is an inhibitory junction potential (IJP) composed of a fast hyperpolarization (fIJP) followed by a slower component (sIJP). fIJPs are mediated by purines (Crist et al. 1992; Gallego et al. 2006; Hwang et al. 2012), and sIJPs are mediated largely by NO (Dalziel et al. 1991; Stark et al. 1991; Keef et al. 1993). Peptide responses can be resolved with higher frequency or stimulation durations of many seconds (Keef et al. 2013). Purine neurotransmitters bind to purinergic receptor sub-type (P2Y1) receptors and activate small conductance Ca^{2+}-activated K^{+} (SK) channels in post-junctional cells, based on studies using receptor antagonists (Gallego et al. 2006; Hwang et al. 2012), SK channel blockers (Banks et al. 1979; Spencer et al. 1998a) and P2ry1−/− mice (Gallego et al. 2012; Hwang et al. 2012).

SMCs have been considered the site of transduction of enteric inhibitory neurotransmission, but recent studies show that interstitial cells labelled by antibodies to platelet-derived growth factor receptor α (PDGFRα+) cells are the post-junctional cells with dominant expression of P2Y1 receptors and SK channels and the cells that generate hyperpolarization in response to P2Y1 agonists (Kurahashi et al. 2011, 2012, 2014; Baker et al. 2013; Peri et al. 2013). PDGFRα+ cells lie in close proximity to varicose nerve terminals in GI smooth muscle tissues (Iino & Nojyo, 2009; Cobine et al. 2011; Kurahashi et al. 2011, 2012; Blair et al. 2012; Grover et al. 2012; Tamada & Hashitani, 2014). Functional data show that PDGFRα+ cells respond to purinergic agonists and generate outward currents, Ca^{2+} transients and hyperpolarization responses consistent with responses of whole muscles to purine neurotransmitters (Kurahashi et al. 2011; Baker et al. 2013; Lee et al. 2013). SMCs, stimulated directly with purine agonists, generate either no response or small inward currents and depolarization. Lastly, electron microscopy has demonstrated gap junctions between PDGFRα+ cells and SMCs in GI muscles (Komuro et al. 1999; Horiguchi and Komuro, 2000; Fujita et al. 2003), suggesting that fIJPs can be conducted through low resistance pathways from PDGFRα+ cells to SMCs. In spite of this evidence, direct innervation of PDGFRα+ cells and an appropriate sequence of activation (i.e. nerve, PDGFRα+ cells, SMC) have not been demonstrated in intact muscles. In this study we used Ca^{2+} indicators loaded into cells within intact muscles and confocal microscopy to investigate the sequence of activation of cells in response to purinergic neurotransmission.

Methods

Animals

PDGFRαtm11(EGFP)Sor/P2ry1−/−, B6.129P2-P2ry1tm1Bhk/J, and SM-eGFP B6.Cg-Tg(Myh11-cre,-EGFP2Mik/B) mice were purchased from the Jackson Laboratory (Bar Harbor, ME, USA). Animals between the ages of 5 and 8 weeks (aged-matched of either sex) were anaesthetized by inhalation of isoflurane (Baxter, Deerfield, IL, USA) and exsanguinated after cervical dislocation before removing the entire GI tracts. The use of animals and experiments performed were in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. The Institutional Animal Use and Care Committee at the University of Nevada approved all procedures.

Tissue preparation

Isolated colons were bathed in Krebs-Ringer bicarbonate solution (KRB) and opened along the mesenteric border. Contents were washed away with KRB. Distal colons (1.5–2.0 cm), 1.5 cm rostral to the anus, were used for these experiments.
Drugs and solutions

Tissues were maintained by constant perfusion with KRB containing (mmol l⁻¹): NaCl, 120.35; KCl, 5.9; NaHCO₃, 15.5; NaH₂PO₄, 1.2; MgCl₂, 1.2; CaCl₂, 2.5; and glucose, 11.5. KRB was bubbled with a mixture of 97% O₂ – 3% CO₂ and warmed to 37 ± 0.2°C. 18-b-Glycyrrhetinic acid, atropine, Nω-nitro-L-arginine (L-NNA), ATP, ADP, β-NAD and octanol were purchased from Sigma-Aldrich (St Louis, MO, USA).

Figure 1. Ca²⁺ transients in PDGFRα⁺ cells in response to exogenous purines
Ca²⁺ transients occurred spontaneously in PDGFRα⁺ cells but were enhanced by ATP (100 μM), ADP (100 μM) and β-NAD (100 μM) (n = 10 for each purine, P = 0.001). D and E, Ca²⁺ transients were also stimulated by the P2Y1 agonist MRS-2365 (1 μM) and UTP (100 μM) (n = 10 and n = 8, respectively; P = 0.001). F–J, Ca²⁺ responses of PDGFRα⁺ cells to purines after pre-treatment with the P2Y1 receptor antagonist, MRS-2500 (1 μM). Responses to ATP (100 μM), ADP (100 μM) and UTP (100 μM) were reduced in the presence of MRS-2500, but not blocked (F, G and J, respectively; n = 6 each, P = 0.001). MRS-2500 abolished Ca²⁺ transient responses to β-NAD (H) and MRS-2365 (J). K, summary of changes in Ca²⁺ transients evoked by purines under control conditions and after addition of MRS-2500. L, summary graph of the effects of TTX (P = 0.008) and MRS-2500 (P = 0.02) on the spontaneous Ca²⁺ transients in PDGFRα⁺ cells in wild-type (WT) muscles. Spontaneous Ca²⁺ transients were also reduced in PDGFRα⁺ cells of P2ry¹⁻/⁻ muscles (n = 5, P = 0.01, raw data traces not shown). Asterisks denote motion/focus artifacts in all panels.
N-Methanocarba-2MeSADP (MRS-2365) and 2-iodo-6-(methylamino)-9H-purin-9-yl] 2 (phosphonoxy) bicyclohexane 1 methanol dihydrogen phosphate ester tetrammonium salt (MRS-2500) were purchased from Tocris Bioscience (Ellisville, MO, USA). All drugs were dissolved in the solvents recommended by the manufacturer to make stock solutions and then dissolved in KRB to the final dilutions desired for experimental tests. 4°C and with secondary antibodies for 1 h at room temperature. The antibodies and dilutions were used as previously described (Kurahashi et al. 2011; Baker et al. 2013). Whole mounts were examined with a Zeiss LSM 510 Meta laser scanning confocal microscope. Confocal micrographs displayed are digital composites of Z-series scans of 0.5–1.0 μm optical sections through a depth of 5–40 μm. Final images were constructed using Zeiss LSM software.

### Immunohistochemistry

Whole mount sections of distal colon were studied with immunohistochemical techniques. Tissues were fixed in acetone or paraformaldehyde (4°C; 10 min) as previously described (Baker et al. 2013) Following fixation, preparations were washed for 30 min in PBS (0.1 M, pH 7.4). Non-specific antibody binding was reduced by incubation in 1% BSA for 1 h at room temperature. The tissues were incubated with primary antibodies for 48 h at 4°C and with secondary antibodies for 1 h at room temperature. The antibodies and dilutions were used as previously described (Kurahashi et al. 2011; Baker et al. 2013). Whole mounts were examined with a Zeiss LSM 510 Meta laser scanning confocal microscope. Confocal micrographs displayed are digital composites of Z-series scans of 0.5–1.0 μm optical sections through a depth of 5–40 μm. Final images were constructed using Zeiss LSM software.

### Fluorescence activated cell sorting (FACS)

Distal colon muscles of PDGFRα<sup>tm11(EGFP)Sor/J</sup> mice and SM-eGFP B6.Cg-Tg<sup>Myh11-cre,EGFP2MiR</sup> J mice were dissected as described above. Muscles were equilibrated in Ca<sup>2+</sup>-free Hanks solution for 30 min and then triturated to disperse cells, as previously described (Baker et al. 2013). PDGFRα<sup>+</sup> cells (enhanced green fluorescent protein (eGFP) in nuclei) and SMC-eGFP cells were sorted by FACS with a Becton-Dickinson FACSAria II instrument.

### Figure 2. Ca<sup>2+</sup> responses of PDGFRα<sup>+</sup> cells to nerve stimulation

A, images showing Ca<sup>2+</sup> responses at different time points during EFS. Arrows show locations of PDGFRα<sup>+</sup> cells, nerve fibres (NF) and SMCs. Scale bar in final panel is 20 μm and pertains to all images. Representative traces of Ca<sup>2+</sup> transients in nerve fibres (NF), PDGFRα<sup>+</sup> cells and SMCs evoked by EFS at different frequencies: responses to single pulse (1 P) (B), 5 Hz (C), 10 Hz (D) and 20 Hz (E) are shown. Stimulus trains were delivered for 1 s (denoted by grey bars below each set of traces and by dotted lines through the traces). Asterisks denote bleedthrough artifacts that usually came from SMCs signals because these cells lay beneath all other cells. In A, Control shows PDGFRα<sup>+</sup> cells clearly distinguishable by eGFP expression in nuclei, EFS (10 Hz) activated nerve bundles (NF) immediately after the onset of EFS (background prior to simulation was subtracted), Ca<sup>2+</sup> responses in PDGFRα<sup>+</sup>-IM were activated soon after responses in neurons (i.e. 0.28 s after onset of EFS (D, G); background and activated nerve bundle images were subtracted). Ca<sup>2+</sup> responses occurred in SMCs subsequent to PDGFRα<sup>+</sup>-IM (B–E). Typically a small decrease in Ca<sup>2+</sup> fluorescence was noted during EFS (i.e. 0.34 s after onset of EFS, D, G) and then an increase in Ca<sup>2+</sup> fluorescence was noted after cessation of EFS (B–F), or in this case about 2 s after initiation of EFS (background and activated nerve bundle images were subtracted). A summary of the latencies (ms) from the start of EFS to the peaks of Ca<sup>2+</sup> transients in PDGFRα<sup>+</sup> cells and SMCs is shown in F (n = 16). A comparison between latencies (ms) from the start of EFS to the initiation of Ca<sup>2+</sup> transients in PDGFRα<sup>+</sup> cells and SMCs at 10 Hz is shown in G (n = 16).
using an excitation laser (488 nm) and emission filter (530/30 nm). Sorting was performed using a 130 μm nozzle at a sheath pressure of 12 p.s.i. (~82.8 kPa) and sort rate of 1000–3000 events s⁻¹. Live cells, gated on exclusion of Hoechst 33258 viability indicator (data not shown), were subsequently gated on eGFP fluorescence intensity.

**RNA extraction and quantitative PCR (qPCR)**

Total RNA was isolated from purified PDGFRα⁺ cells, SMCs and dispersed colonic cells before sorting (i.e. representing the total cell population from the tunica muscularis), using an illustra RNAspin Mini RNA Isolation Kit (GE Healthcare, Piscataway, NJ, USA), and first-strand cDNA was synthesized using SuperScript III (Life Technologies, Grand island, NY, USA), according to the manufacturer’s instructions. The PCR primers used and their GenBank accession numbers are listed in Table 1. Using GoTaq DNA Polymerase (Promega, Madison, WI, USA), PCR products were analysed on 2% agarose gels and visualized by ethidium bromide. qPCR was performed with the same primers as PCR using SYBR green chemistry on the 7500 HT Real-time PCR System (Applied Biosystems, Foster City, CA, USA) and analysed as previously described (Baker et al. 2013).

**Calcium imaging**

Distal colon muscles were pinned to the base of a Sylgard-coated dish. After an equilibration period of 1 h, the preparation was loaded with Oregon Green 488 BAPTA-2 AM (10 μg ml⁻¹; Life Technologies) in a solution of 0.02% DMSO and 0.01% non-toxic detergent Cremophor EL for 30 min at 25°C. After incubation, the preparation was perfused with warmed Krebs solution (37°C for 40 min) to de-esterify the dye.

PDGFRα⁺ cells within circular muscle bundles were identified unequivocally by the eGFP reporter expressed in their nuclei. Ca²⁺ responses to stimulation of intrinsic neurons were measured with a spinning-disc confocal microscope (CSU-X1; spinning disk, Yokogawa Electric, Tokyo, Japan) mounted on an upright Nikon Eclipse FN1 microscope equipped with a 60× lens, Nikon CFI Fluor 60x 1.00 na⁻¹ (Nikon Instruments, New York, USA). The indicator was excited at 488 nm using a laser coupled to a borealis system (ANDOR Technology, Belfast, UK) to increase laser intensity and uniformity. The fluorescence emission (>515 nm) was detected using a high-speed Andor iXon Ultra EMCCD Camera (ANDOR Technology). Image sequences were collected at 33 frames per second using NIS-Elements software (Nikon Instruments). Movies and image sequences of Ca²⁺ activity in PDGFRα⁺ cells were processed and analysed using custom software (Volumetry G8a, G.W.H.).

**Electrical field stimulation (EFS)**

Two parallel platinum electrodes were placed on either side of the colonic muscle sheets described above. Intrinsic neurons were excited by square wave pulses of EFS (one pulse or 5–20 Hz, 0.5 ms pulse durations; 1 s trains) delivered by a Grass S48 stimulator (Quincy, MA, USA). Ca²⁺ and electrophysiological responses induced by EFS were abolished by pretreatment with TTX (1 μM, data not shown). Ca²⁺ imaging and electrophysiological recording were performed in separate experiments. Circular muscle cells were impaled with glass microelectrodes, and transmembrane potentials were measured with a high impedance electrometer (Axoclamp 2B; Axon Instruments/Molecular Devices, Sunnyvale, CA, USA). Membrane potential information was digitized using a Digidata 1322A (Axon Instruments) and recorded by a computer running Axoscope 9.2 software (Axon Instruments).

**Statistical analysis**

Figures displayed were made from digitized data using Adobe Photoshop 4.0.1 (Adobe, Mountain View, CA, USA), Clampfit software (Molecular Devices), Corel Draw 12 (Corel, Ontario, Canada), Excel and PowerPoint 2011 (Microsoft, Redmond, WA, USA). The bar graphs represent the means from each experiment and ‘n’ values refer to the number of animals used for each measurement. Data are expressed as means ± SEM. Statistical significance was calculated using either Student’s t test or a one-way ANOVA followed by a post hoc Newman–Keuls test. P values of < 0.05 were considered to represent significant changes.

Several parameters were calculated from recordings of Ca²⁺ activity, including: (i) durations of responses in nerve fibres, PDGFRα⁺ cells and SMCs; (ii) latency from the initiation of EFS to the peaks of initial Ca²⁺ transients in PDGFRα⁺ cells; and (iii) the latency from the initiation of EFS to the peak of the initial Ca²⁺ transient in SMCs using NIS-Elements software (Nikon Instruments).

Several parameters of electrical activity were also analysed: (i) resting membrane potential; (ii) action potential number before and after EFS; (iii) amplitude of fIJP; (iv) latency from start of EFS to fIJP peak; and (v) latency from the initiation of EFS to the first action potential peak. These parameters were calculated with pCLAMP software (Molecular Devices). The following
Abbreviations are used throughout the analysis and figures (*, cells; **, motion/ focus/ bleed - through artifact). P values are reported in figures as *** ≤ 0.001, ** ≤ 0.01, * ≤ 0.05 and not significant (NS ≥ 0.05).

Results

Ca²⁺ signalling in intramuscular PDGFRα⁺ cells

Ca²⁺ imaging was performed on flat - sheet colonic muscle preparations from PDGFRα⁺/EGFP/Sor/J mice to examine spontaneous Ca²⁺ transients and purinergic responses in PDGFRα⁺ cells. This allowed unequivocal identification of PDGFRα⁺ cells (by EGFP expression in nuclei) and did not obscure resolution of cytoplasmic Ca²⁺ transients. Intramuscular PDGFRα⁺ (PDGFRα⁺/EGFP) cells were found at an average density of 461 ± 16 cells mm⁻² (n = 20; c = 360) and with an average minimum separation between cell bodies of 35.2 ± 2.7 μm (n = 20; c = 360).

Ca²⁺ transients were resolved under basal conditions in several PDGFRα⁺ cells within a given field (spontaneous Ca²⁺ transients occurred in 20.4 ± 3% of cells at an average of 10.2 ± 1.2 events min⁻¹ (range 2–12 events min⁻¹); n = 12). Spontaneous Ca²⁺ transients were not resolved in the remaining PDGFRα⁺ cells. TTX (1 μM) decreased, but did not abolish, spontaneous Ca²⁺ transients in PDGFRα⁺ cells (5.6 ± 0.7 events min⁻¹ after TTX vs. 8.9 ± 0.8 events min⁻¹ in these muscles before TTX; n = 5, P = 0.01; Fig. 1L).

Next we examined responses of PDGFRα⁺ cells to a variety of purine agonists and antagonists. All of these experiments were performed in the presence of l-NNa (100 μM) and atropine (1 μM) to reduce contamination from nitricergic and cholinergic responses. The average occurrence of spontaneous Ca²⁺ transients in PDGFRα⁺ cells before addition of purines was 9.3 ± 1.4 events min⁻¹ (n = 20).

ATP (100 μM) increased the Ca²⁺ transients in PDGFRα⁺ cells (Fig. 1A and K). These responses were typically characterized by an initial sustained rise in fluorescence that tapered off gradually and lasted 3.6 ± 0.42 s (n = 10; Fig. 1A). The sustained rise was followed by oscillatory Ca²⁺ waves that lasted through the recording period (Fig. 1A). ATP increased the Ca²⁺ transients to 23.2 ± 1.5 events min⁻¹ (n = 10, P = 0.001; Fig. 1A and K). ADP (100 μM) evoked Ca²⁺ responses consisting of a sustained Ca²⁺ transient that tapered off gradually and lasted 6.2 ± 0.75 s (n = 10; Fig. 1B). The sustained phase was followed by Ca²⁺ oscillations at 24.4 ± 1.34 events min⁻¹ (n = 10, P = 0.001; Fig. 1B and K).

β-NAD, a candidate purine neurotransmitter in GI muscles (Mutafova-Yambolieva et al. 2007), also evoked Ca²⁺ responses in PDGFRα⁺ cells similar those evoked by ATP and ADP (Fig. 1C). β-NAD (100 μM) increased the activity of Ca²⁺ transients and included an initial sustained rise in Ca²⁺ lasting 2 ± 0.2 s (n = 10; Fig. 1C) and then Ca²⁺ oscillations averaging 12.6 ± 1 events min⁻¹ (n = 10, P = 0.01; Fig. 1C and K). The effects of UTP were also examined. UTP (100 μM) evoked Ca²⁺ responses in the PDGFRα⁺ cells consisting of a sustained Ca²⁺ transient lasting 3.7 ± 0.42 s (n = 8; Fig. 1E). The sustained phase was followed by increased Ca²⁺ oscillations at 16.4 ± 0.84 events min⁻¹ (n = 8, P = 0.001; Fig. 1E and K). Both spontaneously active and quiescent cells responded to exogenous purines. These experiments showed that multiphasic Ca²⁺ responses are elicited in PDGFRα⁺ cells by a variety of naturally occurring purines.

Role of P2Y1 receptors in purinergic responses of PDGFRα⁺ cells

Molecular studies have shown robust expression of P2Y1 receptors in PDGFRα⁺ cells of colon, fundus and bladder.

| Table 1. Summary of gap junction gene primer sequences |
|-----------------------------|-----------------------------|-----------------------------|
| Gene | Primer sequence | GenBank accession number |
| mGapdh-F | GCCGATGCCCCCATGTTGGTA | NM_008084 |
| mGapdh-R | GGGTTGCGATGATGCCATGGGAC | NM_008772 |
| mGja1-F | ACCGAGTGGCTTGTGCGGT | NM_008773 |
| mGja1-R | CCGGCTTTGAGTCGGGCA | NM_020621 |
| mGja5-F | AGACACATCAATGGCACTGGGA | NM_183168 |
| mGja5-R | CAGACAGCTCAAGGCAACACC | NM_027571 |
| mGja7-F | GGGCCTCAATGCCCCAACCC | NM_020621 |
| mGja7-R | GCCATTGCAAAGGCGCAGT | NM_183168 |
| mGb1-F | ACCGACCTGTGTCGTACGAC | NM_027571 |
| mGb1-R | CGCGGGGCATGCGACAATA | NM_027571 |
| mGb2-F | GGCCTGACGTCACTGAAACGCCTG | NM_027571 |
| mGb2-R | TCTTTCGGCTTTGTTCCACC | NM_027571 |

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muscles (Kurahashi et al. 2011; Baker et al. 2013; Lee et al. 2013). Previous studies have shown that purinergic stimulation activates outward currents in single isolated PDGFRα⁺ cells (Kurahashi et al. 2011). Therefore, we evaluated the role of P2Y1 receptors in mediating Ca²⁺ responses to purines in PDGFRα⁺ cells in situ.

Similar to responses to biological purines, a robust increase in Ca²⁺ transients was elicited in PDGFRα⁺ cells by the P2Y1 receptor agonist MRS-2365 (1 μM; Fig. 1D). MRS-2365 increased Ca²⁺ transients to 30.5 ± 1.2 events min⁻¹ (n = 10; P = 0.001; Fig. 1D and K). These responses were also characterized by an initial sustained rise in Ca²⁺ that lasted 8.2 ± 0.58 s (n = 10; Fig. 1D).

Pretreatment of muscles with MRS-2500 (1 μM), a highly selective antagonist of P2Y1 receptors, decreased, but did not abolish, spontaneous Ca²⁺ transients in PDGFRα⁺ cells (5.2 ± 1.1 events min⁻¹; n = 5, P = 0.02; Fig. 1L). MRS-2500 abolished Ca²⁺ responses evoked by β-NAD and MRS-2365 (Fig. 1H and I, respectively). Responses to ATP and ADP were attenuated by MRS-2500, and the sustained Ca²⁺ oscillations typical of ATP responses were reduced to 5.1 ± 1.52 events min⁻¹ (n = 6, P = 0.001; Fig. 1F and K). Responses to ADP after MRS-2500 pretreatment were reduced to 2.9 ± 1.3 events min⁻¹ (n = 6, P = 0.001; Fig. 1G and K).

In contrast, MRS-2500 reduced the UTP response by only 58.4 ± 4% (to 6.8 ± 1.3 events min⁻¹, n = 6, P = 0.001; Fig. 1J and K).

**PDGFRα⁺ cell responses to nerve stimulation**

Ca²⁺ responses to EFS (single pulse and 5–20 Hz; 1 s trains; 0.5 ms pulse duration) were characterized in PDGFRα⁺IM cells identified by eGFP in nuclei (Fig. 2A–E). In the presence of L-NNA (100 μM) and atropine (1 μM), EFS triggered rapid transient Ca²⁺ responses in nerve fibres that peaked 60 ± 3.4 ms after initiation of EFS and decayed within 1991 ± 100.69 ms (10 Hz EFS; n = 16; Fig. 2D). Ca²⁺ transients were stimulated in both spontaneously active and quiescent PDGFRα⁺ cells after initiation of EFS, but these responses were delayed slightly relative to the responses in nerve fibres. Ca²⁺ transients were initiated in PDGFRα⁺ cells with a latency of 280 ± 17.4 ms and peaked at 356 ± 16.4 ms after the onset of EFS. Ca²⁺ responses in PDGFRα⁺ cells lasted 1615 ± 94.6 ms at 10 Hz EFS (n = 16; Fig. 2F, G and Supporting Information, Movie S1).

SMCs responded to EFS initially with a decrease in Ca²⁺ fluorescence that began after the activation of Ca²⁺ transients in PDGFRα⁺ cells. Initiation of this drop in fluorescence occurred close in time to the peak of the Ca²⁺ transient in PDGFRα⁺ cells (i.e. latency was 344 ± 15.8 ms at 10 Hz; n = 11; Fig. 3D and G). A majority of SMCs (69%) displayed a decrease in Ca²⁺ fluorescence that lasted 1683 ± 84.3 ms and decreased 16.7 ± 2.3% below basal Ca²⁺ in response to 10 Hz EFS (n = 11, Figs 2D and 3D). A decrease in basal Ca²⁺ could not be resolved in the remainder of the SMCs during EFS. In cells in which Ca²⁺ was reduced, the reduction was sustained during EFS, and then a post-stimulus (rebound) excitation developed and triggered positive Ca²⁺ transients in SMCs. The post-stimulus Ca²⁺ responses in SMCs occurred well after the Ca²⁺ transients in neuronal and PDGFRα⁺ cells had settled and after cessation of EFS. The latency from initiation of EFS to the peak of Ca²⁺ transients in SMCs was 1941 ± 18.2 ms at 10 Hz EFS (n = 16; Fig. 2D and F). The post-stimulus Ca²⁺ transients in SMCs lasted for 804 ± 106.4 ms at 10 Hz EFS (n = 16; Fig. 2D).

These experiments demonstrate a temporal sequence in the responses to enteric motor nerve stimulation: nerve bundles were activated immediately after the onset of EFS, PDGFRα⁺ cells responded after a short latency, and this response was followed rapidly by a dip in Ca²⁺ in SMCs (see time sequence images in Fig. 2A). Figure 2F and G summarizes the latencies after initiation of EFS to Ca²⁺ responses in PDGFRα⁺ cells and SMCs. The appearance of Ca²⁺ transients in PDGFRα⁺ cells prior to SMCs indicates the sequence of activation in response to the purinergic component of neurotransmission.

**Ca²⁺ responses of PDGFRα⁺ cells are mediated via P2Y1 receptors**

Electrophysiological experiments have shown that purinergic neurotransmission is mediated by post-junctional P2Y1 receptors in the murine colon (Gallego et al. 2012; Hwang et al. 2012). Therefore, we examined the effects of a selective P2Y1 antagonist on the Ca²⁺ transients elicited by EFS. Time sequence images of Ca²⁺ transients evoked by EFS show that the responses of PDGFRα⁺ cells and SMCs were inhibited by MRS-2500 (1 μM; Fig. 3A and B). In the presence of MRS-2500, EFS (10 Hz) evoked Ca²⁺ responses in nerve fibres with an average duration of 1727 ± 43.9 ms in comparison to control (1921 ± 70.34 ms; P = 0.03, n = 8; Fig. 3C–F), but Ca²⁺ transients were not resolved in PDGFRα⁺ cells or SMCs in the presence of MRS-2500 (Fig. 3E and F). MRS-2500 also abolished the reduction in Ca²⁺ levels during EFS in SMCs. These results suggest that post-junctional Ca²⁺ transients in PDGFRα⁺ cells and SMCs evoked by EFS were mediated by P2Y1 receptors.

**Lack of Ca²⁺ responses to EFS in PDGFRα⁺ cells and SMCs from P2ry1⁻/⁻ mice**

Purinergic IJPs were shown previously to be absent in P2ry1⁻/⁻ mice (Gallego et al. 2012; Hwang et al.
PDGFRα+ cells of P2ry1+/− colonic muscles displayed fewer spontaneous Ca2+ transients than observed in cells of P2ry1+/+ muscles (i.e. 16.2 ± 2.1% of spontaneous Ca2+ transients in P2ry1+/− cells occurred at an average of 4.8 ± 0.66 events min⁻¹, P = 0.01; n = 6; Fig. 1L).

EFS of muscles from P2ry1−/− mice elicited Ca2+ transients in nerve fibres with an average duration of 1766 ± 33.4 ms (n = 6; Fig. 4B and Movie S2) but failed to evoke Ca2+ responses in PDGFRα+ cells or SMCs (Fig. 4B and C). Time sequence images also demonstrated the lack of Ca2+ responses in PDGFRα+ cells and SMCs in P2ry1−/− colonic muscles (Fig. 4C), thus confirming the importance of P2Y1 receptors in mediating post-junctional Ca2+ transients in PDGFRα+ cells and SMCs in purinergic neurotransmission.
**Importance of P2Y1 receptors in IJPs**

Intracellular microelectrode recordings were also performed in separate experiments to relate the temporal sequence of \(Ca^{2+}\) transients to the well-characterized electrophysiological response to purinergic neurotransmission. In the presence of L-NNA and atropine, EFS (one pulse and 5–20 Hz) evoked IJPs followed by post-stimulus depolarization responses, as previously described in several species (Fig. 5A and C). IJPs averaged 29.5 ± 2 mV in amplitude (\(n = 6\); Fig. 5A and D) and 1832 ± 45.8 ms in duration in response to 10 Hz EFS (\(n = 6\); Fig. 6B). Pretreatment of muscles with MRS-2500 had no initial effect on membrane potential (i.e. \(-56 ± 1.7\) mV in control vs. \(-57 ± 2.2\) mV after addition of MRS-2500; \(n = 6\); Fig. 5A and B), but this compound blocked fIJPs at all frequencies below 20 Hz. A small component of hyperpolarization escaped block by MRS-2500 at 20 Hz (4.5 ± 1.3 mV, \(n = 6\); Fig. 5B and D). A train of action potentials followed cessation of EFS (post-stimulation excitation), which has been described previously in GI muscles (Bennett, 1966; Wood & Brann, 1986; Ward *et al.* 1992) (Fig. 5A). For example, before EFS the spontaneous action potential occurred at 3 ± 0.1 events 10 s\(^{-1}\), and this was increased significantly after EFS (10 Hz) to 6.2 ± 0.6 events 10 s\(^{-1}\) (\(P = 0.002, n = 6\); Fig. 5A and C). Post-stimulus excitation was also blocked by MRS-2500 and action potentials after EFS did not exceed the rate of spontaneous action potential generation (2.9 ± 0.08 events 10 s\(^{-1}\) at 10 Hz EFS, \(n = 6\); Fig. 5C).

**Figure 4. Ca\(^{2+}\) responses in PDGFR\(\alpha^+\) cells and SMCs evoked by EFS were absent in P2ry1\(^{−/−}\) mice**

A, immunolabeling of whole mounts of colon from P2ry1\(^{−/−}\) mice with PDGFR\(\alpha\) (red, Aa) and SK3 (red, Ad) antibodies showed double labelling of PDGFR\(\alpha^+\) cells (green eGFP nuclei, Ab, Ae; merged images are in Ac and Af). Scale bar in Af is 50 \(\mu\)m and pertains to all panels in A. Ca\(^{2+}\) transients evoked by EFS were absent in PDGFR\(\alpha^+\) cells and SMCs in muscles of P2ry1\(^{−/−}\) mice (B). The stimulus train (1 s) is denoted by the red bar and dotted line through the traces. Time sequence images demonstrate the absence of Ca\(^{2+}\) transients in PDGFR\(\alpha^+\) cells and SMCs in muscles of P2ry1\(^{−/−}\) mice (C). White arrowheads indicate PDGFR\(\alpha^+\) cells or nerve fibres. Scale bar in C (right panel) is 20 \(\mu\)m and pertains to all panels in C.
The latencies from the onset of EFS (5–20 Hz) to the peaks of fIJPs and the peaks of the Ca\(^{2+}\) transients in PDGFR\(\alpha\)+ cells were highly correlated \((R^2 = 0.99; \text{Fig. } 6 \text{C})\). A strong correlation was also observed between the latency of the first post-stimulus action potential and the Ca\(^{2+}\) transients observed in SMCs after cessation of EFS \((R^2 = 0.96; \text{Fig. } 6 \text{D})\). Taken together, these data demonstrate a consistent relationship between Ca\(^{2+}\) transients in PDGFR\(\alpha\)+ cells and electrophysiological events in response to stimulation of motor neurons and show that PDGFR\(\alpha\)+ cells are likely to mediate fIJPs in colonic muscles.

**Expression of gap junctions in PDGFR\(\alpha\)+ cells**

In order for responses developed in PDGFR\(\alpha\)+ cells to conduct to SMCs, some form of low resistance pathway(s) must connect these cells. Morphological studies have shown that PDGFR\(\alpha\)+ cells are coupled to SMCs via gap junctions (Komuro et al. 1999; Horiguchi & Komuro, 2000; Fujita et al. 2003; Iino et al. 2009), but the nature and composition of gap junctions that form between these cells has not been investigated. Connexin (Cx) 40, 43 and 45 have been reported to be expressed in colonic muscles (Li et al. 1993; Mikkelsen et al. 1993; Nakamura et al. 1998; Seki & Komuro, 2001; Wang & Daniel, 2001), and more recently, human colon was shown to express Cx 32 and 26 (Kanczuga-Koda et al. 2004). We characterized the expression of gap junction genes in PDGFR\(\alpha\)+ cells and SMCs using qPCR on extracts of cells purified by FACS and found higher expression of Gjb1 (Cx 32) and Gjb2 (Cx 26) in PDGFR\(\alpha\)+ cells in comparison to unsorted cells (fold change = 2.8 ± 0.16, \(P = 0.0004; \text{fold change} = 2.1 ± 0.12, \(P = 0.001\), respectively; Fig. 7A). We also observed higher expression of Gja5 (Cx 40) in PDGFR\(\alpha\)+ cells (fold change = 1.8 ± 0.12, \(P = 0.003\); Fig. 7A). Gja7 (Cx 45) expression was not significantly

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**Figure 5. Electrical responses (IJPs) to EFS**

*Panel A*, electrical activity recorded from intact colonic muscles in the presence of L-NNA (100 \(\mu\)M) and atropine (1 \(\mu\)M). EFS single pulse (1 P) and 5–20 Hz elicited IJPs, followed by post-stimulus excitation consisting of a train of action potentials (AP). Previous studies have described these responses as purinergic fast IJPs (fIJPs). *Panel B*, MRS-2500 (1 \(\mu\)M) abolished fIJPs evoked by 1 P, 5 and 10 Hz stimuli. A small component persisted after MRS-2500 at 20 Hz (4.5 ± 1.2 mV, \(n = 6\)). *Panel C*, summary of the average number of post-stimulus action potentials in controls and in the presence of MRS-2500 at different frequencies (i.e. one pulse and 5–20 Hz; each bar in the graph represents the average number of action potentials within 10 s in response to EFS). Note that the post-stimulus activation of action potentials was also inhibited by MRS-2500 (\(n = 6\); each bar in the graph represents the average IJP amplitude).

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increased in PDGFRα+ cells (fold change = 1.1 ± 0.07, P = 0.16; Fig. 7A). Gja1 (Cx 43) showed lower expression in PDGFRα+ cells in comparison to the population of unsorted cells (fold change = −4.5 ± 0.01, P = 0.001; Fig. 7B).

Sorted SMCs showed higher expression of Gja7 (Cx 45) in comparison to unsorted cells (fold change = 1.4 ± 0.15, P = 0.008; Fig. 7B) and significantly lower expression of other gap junctions: Gja1 (Cx 43) compared to unsorted cells (fold change = −4.2 ± 0.4, P = 0.0003; Fig. 7B); Gja5 (Cx 40) compared to unsorted cells (fold change = −1.6 ± 0.16, P = 0.0001; Fig. 7B); and Gjb1 (Cx 32) expression (fold change = −2.9 ± 0.2, P = 0.0001; Fig. 7B). And a reduction in Gjb2 (Cx 26) expression was also observed (fold change = −2 ± 0.14, P = 0.0001; Fig. 7B). It was also noted that the highest gap junction levels in PDGFRα+ cells and SMCs were Gja1 (Cx 43) and Gja7 (Cx 45), regardless of the ratio of expression to unsorted cells. These results suggest that PDGFRα+ cells and SMCs express several gap junction genes that may be involved in electrical coupling between these cells.

**Gap junction blocking drugs blocked purinergic Ca\(^{2+}\) responses in SMCs**

We examined the role of gap junctions in mediating the purinergic signalling in post-junctional cells with the gap junction inhibitors 18-β-glycyrrhetinic acid (18-β-GA) and octanol. Pretreatment of colonic muscles with 18-β-GA (40 μM; for 10–12 min in the presence of L-NNA and atropine) did not affect Ca\(^{2+}\) responses to EFS (10 Hz) in nerve fibres (duration of 1907 ± 83.2 ms, P = 0.54, n = 6; Fig. 8C) or PDGFRα+ cells (duration of 1624 ± 106.6 ms, P = 0.85; latency from EFS 332 ± 29.6 ms, P = 0.45, n = 6; Fig. 8C and D). In contrast, Ca\(^{2+}\) responses in SMCs...
were reduced in duration (to 483 ± 67.2 ms, P = 0.01, n = 6; Fig. 8C) but no change in the latency to the peak of post-stimulus Ca\(^{2+}\) transients in SMCs was observed (latency from EFS 1938 ± 94.2 ms, P = 0.61, n = 6; Fig. 8D). Higher concentrations of 18-β-GA (100 μM) blocked all Ca\(^{2+}\) responses to EFS in SMCs but had no effect on the duration of Ca\(^{2+}\) transients in nerve fibres (1809 ± 70.8 ms, P = 0.12, n = 6; Fig. 8B and C) or PDGFRα\(^+\) cells (1475 ± 83.8 ms, P = 0.21, n = 6; Fig. 8B and C) and no significant effects were noted in the latency between onset of EFS and the peaks of Ca\(^{2+}\) transients in PDGFRα\(^+\) cells (318 ± 24.9 ms, P = 0.26, n = 6; Fig. 8D). Higher concentrations of 18-β-GA (100 μM) blocked all Ca\(^{2+}\) responses to EFS in SMCs but had no effect on the duration of Ca\(^{2+}\) transients in nerve fibres (1809 ± 70.8 ms, P = 0.12, n = 6; Fig. 8B and C) or PDGFRα\(^+\) cells (1475 ± 83.8 ms, P = 0.21, n = 6; Fig. 8B and C) and no significant effects were noted in the latency between onset of EFS and the peaks of Ca\(^{2+}\) transients in PDGFRα\(^+\) cells (318 ± 24.9 ms, P = 0.26, n = 6; Fig. 8D).

A second gap junction blocking drug, octanol, was tested on post-junctional responses to EFS. Octanol (300 μM) caused a small, but significant, reduction in the Ca\(^{2+}\) responses evoked in nerve fibres by EFS (10 Hz) (from a control duration of 2009 ± 44.4 ms to 1867 ± 42.7 ms in the presence of octanol; P = 0.04, n = 6; Fig. 9C) and a reduction in PDGFRα\(^+\) cells (reduced from a control duration of 1671 ± 101.76 ms to 1276 ± 65.98 ms; P = 0.008, n = 6; Fig. 9C). Responses of SMCs to EFS were also reduced by octanol (300 μM) (from a control duration of 897 ± 81.9 ms to 537 ± 60.6 ms; P = 0.005, n = 6; Fig. 9C). Octanol (300 μM) had no significant effect on the latencies between the onset of EFS and peak Ca\(^{2+}\) transients in PDGFRα\(^+\) cells (404.3 ± 29.9 ms, P = 0.46, n = 6; Fig. 9D). While octanol appeared to have some non-specific effects on pre-junctional responses in enteric

**Figure 7. Expression of gap junction transcripts in PDGFRα\(^+\) cells**

The relative expression of gap junction gene transcripts (Gja1, Gja5, Gja7, Gjb1, Gjb2) was compared in sorted PDGFRα\(^+\) cells (A), sorted SMCs (B) and unsorted cells (i.e. mixed cell population after enzymatic dispersions of distal colon muscles) by qPCR. A, PDGFRα\(^+\) cell expression of Cx 40 (Gja5), Cx 32 (Gjb1) and Cx 26 (Gjb2) were higher in PDGFRα\(^+\) cells in comparison to other cell types, although the highest transcript levels in PDGFRα\(^+\) cells were Cx 43 (Gja1) and Cx 45 (Gja7). B, transcript expression in sorted SMCs. Cx 45 (Gja7) and Cx 43 (Gja1) were also the most highly expressed gap junction genes in SMCs in comparison to other gap junction genes. The relative expression of each gene was normalized to the housekeeping gene, Gapdh.
neurons, its effects on post-junctional cells were consistent with the effects of 18-β-GA.

**Gap junction blocking drugs inhibited the conduction of purinergic electrical responses to SMCs**

We also recorded electrophysiological responses of colonic muscles to EFS before and after addition of gap junction blockers. 18-β-GA (40 μM) caused a slight depolarization of cells that did not reach statistical significance (from −53 ± 2.2 to −49.5 ± 2.5 mV within 10–12 min after adding 18-β-GA; \( P = 0.29, n = 6 \); Fig. 10A). In the presence of 18-β-GA (40 μM) EFS (5, 10 and 20 Hz) evoked IJPs that were significantly reduced in amplitude (e.g. IJPs at 10 Hz were 29.6 ± 0.8 mV in amplitude under control conditions and 26.9 ± 0.72 mV after 18-β-GA; \( P = 0.01, n = 6 \); Fig. 10C). No significant change was noted in IJP amplitudes with one pulse of EFS (24.7 ± 1 mV, \( P = 0.25 \)). 18-β-GA (40 μM) had no significant effect on the latency of IJP parameters. For example, at 10 Hz EFS the latency between EFS and fIJP was 401 ± 16 ms compared to control value of 363 ± 16.7 ms (\( P = 0.14, n = 6 \)) and the latency between EFS and rebound peak was 1963 ± 53.4 ms compared to control value of 1854 ± 63.7 ms (\( P = 0.22, n = 6 \)). However, in the presence of 18-β-GA (40 μM) post-stimulus action potentials were decreased at all frequencies of EFS tested (i.e. at 10 Hz action potentials were 4.7 ± 0.46 events 10 s\(^{-1}\) compared to control value of 6.4 ± 0.44 events 10 s\(^{-1}\), \( n = 6, P = 0.02; \) Fig. 10E). Pretreatment of colonic muscles with 18-β-GA (100 μM) abolished fIJPs and post-stimulus action potentials (Fig. 10A). Membrane depolarization was also observed with this concentration of 18-β-GA (i.e. to −44 ± 1.6 mV, \( P = 0.001 \)).

Octanol (300 μM) caused slight hyperpolarization, but this effect did not reach statistical significance (−58 ± 1.3 mV compared to control value of −55 ± 1.7 mV, \( P = 0.16, n = 6 \); Fig. 10B). In the presence of octanol (300 μM) IJPs were reduced in amplitude at all frequencies tested (e.g. 18.7 ± 3.1 mV at 10 Hz EFS.

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**Figure 8.** 18-β-glycyrrhetinic acid (18-β-GA) failed to block EFS-evoked responses in PDGFRα+ cells but blocked responses in SMCs

A, an example of Ca\(^{2+}\) transients activated by EFS (10 Hz) under control conditions (l-NNA and atropine present, \( n = 6 \)). The stimulus train (1 s) is denoted by the grey box and dotted lines through the trace. EFS evoked Ca\(^{2+}\) transients in PDGFRα+ cells with a short latency after onset of EFS and in SMCs after cessation of stimulation. 18-β-GA (100 μM) blocked Ca\(^{2+}\) responses evoked by EFS in SMCs but did not affect the responses in PDGFRα+ cells significantly (B). C, summary of the duration of Ca\(^{2+}\) transients in nerve fibre bundles (black bars), PDGFRα+ cells (white bars) and SMCs (grey bars) before and after addition of 18-β-GA (40 and 100 μM). D, summary graph of the average latency in Ca\(^{2+}\) responses in PDGFRα+ cells and SMCs after the onset of EFS to the first Ca\(^{2+}\) transient peak before and after the addition of 18-β-GA (40 and 100 μM). Little change was noted in the latencies in nerve fibres and PDGFRα+ cells after 18-β-GA (100 μM), but responses were blocked in SMCs in the presence of 100 μM 18-β-GA.
compared to control value of 31.5 ± 1.8 mV, P = 0.005, n = 6; Fig. 10D). Octanol (300 μM) had no significant effect on the latency of IJPs. For example, at 10 Hz EFS the latency between EFS and fIJPs averaged 397 ± 45.1 ms, compared to control value of 376 ± 28.3 ms (P = 0.71, n = 6), and the latency between EFS and the post stimulus rebound response was 1894 ± 59.7 ms, compared to control value of 1783 ± 67.1 ms (P = 0.24, n = 6). In the presence of octanol (300 μM) there was a marked decrease in post-stimulus action potentials after cessation of EFS at all frequencies tested (e.g. at 10 Hz EFS action potentials averaged 4.2 ± 0.32 events 10 s⁻¹ compared to control value of 7.1 ± 0.4 events 10 s⁻¹; n = 6, P = 0.0002; Fig. 10F). Pretreatment of muscles with higher concentrations of octanol (700 μM) abolished fIJPs and post-stimulus action potentials (Fig. 10B), and it should be noted that hyperpolarization was also observed with this concentration of octanol (−63 ± 1.9 mV, P = 0.01; Fig. 10B).

Discussion

This study provides evidence that PDGFRα⁺-IM cells in colonic muscles are innervated by enteric inhibitory motor neurons. PDGFRα⁺-IM cells were responsive to exogenous purines and the responses were blocked by MRS-2500. With cholinergic and nitric components of responses blocked, EFS evoked Ca²⁺ transients in post-junctional cells (PDGFRα⁺-IM cells and SMCs). Post-junctional responses were abolished by MRS-2500 and not observed in muscles with genetic deactivation of P2Y1 receptors. Electrical responses (IJPs) showed similar time courses to the Ca²⁺ transients elicited by EFS. Drugs known to inhibit gap junctions blocked post-junctional responses to EFS in SMCs, but did not block responses in intramuscular nerve processes or PDGFRα⁺-IM cells. These data support the hypothesis that purinergic neurotransmission is transduced by PDGFRα⁺-IM cells and conducts to SMCs via gap junctions. The study shows the first direct evidence for serial activation of post-junctional cells during purinergic neurotransmission.

PDGFRα⁺ cells are abundant in the distal colon, and the cells within muscle bundles (PDGFRα⁺-IM cells) lie in close proximity to varicose processes of motor neurons (Kurahashi et al. 2011, 2012; Blair et al. 2012). This morphology occurs throughout various regions of the GI tracts of all mammalian species studied, including humans, as determined by light and electron microscopy (Komuro et al. 1999; Fujita et al. 2003; Iino et al. 2009; Grover et al. 2012). PDGFRα⁺ cells
also possess the molecular apparatus for mediating purinergic signalling. Exogenous purines elicit Ca\(^{2+}\) transients in PDGFR\(\alpha\) cells of the gastric fundus, and cells from the fundus and colon express P2Y1 receptors and SK3 channels that are the basis for purinergic IJPs (Mutafova-Yambolieva et al. 2007; Kurahashi et al. 2011; Baker et al. 2013; Peri et al. 2013). The present study unites morphological and physiological observations by showing that PDGFR\(\alpha\)-IM cells are innervated, respond directly to neurotransmitters released from enteric motor neurons and conduct responses to SMCs.

PDGFR\(\alpha\)-IM cells displayed Ca\(^{2+}\) transients spontaneously and these events were attenuated, but not blocked, by TTX. This suggests that PDGFR\(\alpha\) cells contribute to setting resting smooth muscle excitability through ongoing inhibitory neurotransmission and by intrinsic generation of spontaneous transients outward currents (STOCs). This activity is likely to contribute to tonic inhibition, a basic behaviour in GI motility (Wood, 1972; Waterman & Costa, 1994; Spencer et al. 1998a). PDGFR\(\alpha\) cells have robust expression of SK3 channels that are activated by cytoplasmic Ca\(^{2+}\) and blocked by apamin (Kurahashi et al. 2011). The spontaneous Ca\(^{2+}\) transients in PDGFR\(\alpha\) cells were increased by exogenous purines. Ca\(^{2+}\) transients occur in a stochastic manner, and are dependent upon release of Ca\(^{2+}\) from internal stores via inositol trisphosphate (IP\(_3\)) and ryanodine receptors (Baker et al. 2013; Tamada & Hashitani, 2014). The Ca\(^{2+}\) transients are asynchronous cell to cell, and are likely to cause STOCs in single PDGFR\(\alpha\) cells and spontaneous transient hyperpolarizations (STHs) in intact muscles (Kito et al. 2014). Ca\(^{2+}\) transients in PDGFR\(\alpha\)-IM
cells appear to mediate the spontaneous IJPs in colonic muscles, reported previously, as these events were also sensitive to TTX and apamin (Spencer et al. 1998b; Gil et al. 2010). Summation of spontaneous IJPs and intrinsic STOCs in PDGFRα−IM cells would provide a net hyperpolarizing influence on the smooth muscle and temper basal contractile activity (i.e. a source of tonic inhibition).

P2Y1 receptors mediate purinergic neurotransmission in the gut (Giaroni et al. 2002; Gallego et al. 2006; Grasa et al. 2009; Zhang et al. 2010). Binding of purines to P2Y1 receptors couples to activation of SK channels and generation of fIJPs (Gallego et al. 2006; Grasa et al. 2009; Hwang et al. 2012). We confirmed that fIJPs are blocked by the selective P2Y1 blocker MRS-2500 and also demonstrated that Ca2+ transients activated by neuro-transmitters were blocked by MRS-2500 and absent in P2ryl1−/− mice. Note that Ca2+ transients activated by exogenous ATP, ADP and UTP were attenuated but not fully inhibited by MRS-2500, suggesting that PDGFRα+ cells express purine receptors in addition to P2Y1. P2Y2 receptors are equally sensitive to ATP and UTP (Velazquez et al. 2000), and expression of P2Y2 receptors by PDGFRα+ cells has been reported (Peri et al. 2013). MRS-2500 attenuated half of the UTP responses in the PDGFRα+ cells and this antagonism may attribute to a cross-talk between P2Y2 and P2Y1 receptors through phospholipase C (PLC) pathways (Werry et al. 2003; Baranska et al. 2004). The effects of ADP remaining after MRS-2500 might be mediated by P2Y12 receptors that are also expressed in PDGFRα+ cells (Peri et al. 2013). Binding of P2Y12 receptors might enhance Ca2+ mobilization through IP3 receptors by inhibition of protein kinase A (van der Meijden et al. 2008).

β-NAD, a purinergic neurotransmitter candidate, also elicited Ca2+ responses in PDGFRα− cells, and these responses were blocked by MRS-2500 (Mutafova-Yambolieva et al. 2007; Hwang et al. 2012). A recent study demonstrated differential expression of enzymes involved in purine catabolism in cells within the tunica muscularis (Peri et al. 2013). For example, ectoenzymes Cd38 and Enpp1 are highly expressed in PDGFRα+ cells in comparison to SMCs and interstitial cells of Cajal (ICC). Cd38 is a primary enzyme capable of NAD+ hydrolysis (De Flora et al. 2004). Thus, close apposition of neurotransmitter release sites to PDGFRα+ cells and focalization of mechanisms to deactivate neurotransmitters close to sites of release may limit the post-junctional volume in which effective neurotransmitter concentrations are achieved.

This study measured responses and latencies of the responses of PDGFRα− cells and SMCs to EFS of intrinsic motor neurons. Cholinergic and nitricergic neurotransmission were blocked to focus on responses to purinergic neurotransmission. Ca2+ transients were increased in PDGFRα+ cells by purinergic neurotransmission, and this response slightly preceded the occurrence of an fIJP. Such a Ca2+ response is consistent with activation of SK channels, which are responsible for fIJPs. Shortly after initiation of Ca2+ transients in PDGFRα+ cells, basal Ca2+ was reduced in SMCs. With the resolution of Ca2+ events in this study, an initial rise in Ca2+ was never evoked in SMCs by EFS. SMCs have low levels of SK channel expression and much lower current density attributable to SK channels than PDGFRα+ cells (Mutafova-Yambolieva et al. 2007; Kurahashi et al. 2011; Peri et al. 2013), but without a rise in Ca2+ in SMCs with kinetics similar to the kinetics of the fIJP, there is no mechanism to activate the SK channels that might be available in SMCs. The sustained drop in Ca2+ observed in SMCs during purinergic nerve stimulation is consistent with the idea that hyperpolarization responses developed in PDGFRα+ cells conducted to SMCs and reduced Ca2+ influx by reducing openings of voltage-dependent Ca2+ channels. Decreased Ca2+ in SMCs would be expected to cause relaxation, which is the well-known response to purinergic inhibitory neurotransmission in GI muscles.

Clearly electrical connectivity between PDGFRα+ cells and SMCs must exist if activation of SK3 channels in PDGFRα+ cells is capable of eliciting hyperpolarization in SMCs. Ultrastructural studies have reported gap junctions between PDGFRα+ cells and SMCs (Komuro et al. 1999; Fujita et al. 2003), and multiple gap junction proteins and transcripts, including Cx 40, 43, 45, 26 and 32, are expressed in GI smooth muscle tissues (Li et al. 1993; Mikkelsen et al. 1993; Nakamura et al. 1998; Seki & Komuro, 2001; Wang & Daniel, 2001; Kanczuga-Koda et al. 2004). All these studies reported results from whole muscles in which many cell types exist. We found a number of gap junction genes expressed in PDGFRα+ cells and SMCs. The variety of gap junctions possible between PDGFRα+ cells and SMCs from the genes expressed would provide a source of tonic inhibition and would allow effective neurotransmitter concentration to be maintained.

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The fact that responses were retained in nerves and PDGFRα+ cells in the presence of gap junction inhibitors is consistent with the following concept of purinergic neurotransmission: propagation of electrical signals in neurons, Ca2+ influx into nerve processes, release of neurotransmitter, binding of P2Y1 receptors expressed by PDGFRα+ cells and activation of Ca2+ release in these cells. Gap junctions are not involved in any of these steps, and therefore gap junction blockers did not block these events. Our findings support the hypothesis that post-junctional responses (IJPs) develop in PDGFRα+ cells and conduct to SMCs via gap junctions. Thus, responses in SMCs do not appear to result from direct binding of smooth muscle receptors by purine neurotransmitters.

In summary, this study demonstrates direct functional innervation of PDGFRα+ cells by purinergic enteric inhibitory neurons in colonic muscles. Morphological studies predicted there might be communications between motor neurons and PDGFRα+ cells, but this is the first report showing functional innervation of these cells. Ca2+ transients activated in PDGFRα+ cells are a requisite for the activation of SK channels, a signature of purinergic neurotransmission in GI muscles. Ca2+ transients were observed to occur spontaneously, which might provide part of the ‘tonic inhibition’ imposed on colonic muscles to help maintain the phasic nature of contractions. The number and amplitude of Ca2+ transients were increased by exogenous purines and by purinergic neurotransmitter(s) released from inhibitory motor neurons. This study supports targeted neurotransmission, specialization of post-junctional cells in transducing inputs from motor neurons in GI muscles and a primary role for PDGFRα+ cells in mediating enteric inhibitory neurotransmission in the colon.

References


**Additional information**

**Competing interests**

None

**Author contributions**

Conception and design of the experiments: S.A.B., K.M.S. Collection, analysis and interpretation of data: S.A.B., G.W.H., S.M.W., K.M.S. Drafting the article or revising it critically for important intellectual content: S.A.B., G.W.H., S.M.W., K.M.S. All authors read and approved the manuscript for submission.

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**Supporting information**

The following supporting information is available in the online version of this article.

*Movie S1.* PDGFR\(\alpha^+\) cell Ca\(^{2+}\) responses to nerve stimulation.

*Movie S2.* Ca\(^{2+}\) transients of PDGFR\(\alpha^+\) cells are absent in *P2ry1\(^{−/−}\)* mice.