Thrombin-Mediated Direct Activation of Proteinase-Activated Receptor-2: Another Target for Thrombin Signaling

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ABSTRACT

Thrombin is known to signal to cells by cleaving/activating a G-protein–coupled family of proteinase-activated receptors (PARs). The signaling mechanism involves the proteolytic unmasking of an N-terminal receptor sequence that acts as a tethered receptor-activating ligand. To date, the recognized targets of thrombin cleavage and activation for signaling are PAR1 and PAR4, in which thrombin cleaves at a conserved target arginine to reveal a tethered ligand. PAR2, which like PAR1 is also cleaved at an N-terminal arginine to unmask its tethered ligand, is generally regarded as a target for trypsin but not for thrombin signaling. We now show that thrombin, at concentrations that can be achieved at sites of acute injury or in a tumor microenvironment, can directly activate PAR2 vaso-relaxation and signaling, stimulating calcium and mitogen-activated protein kinase responses along with triggering β-arrestin recruitment. Thus, PAR2 can be added alongside PAR1 and PAR4 to the targets, whereby thrombin can affect tissue function.

Introduction

The cloning of the G-protein–coupled receptor for thrombin, which is responsible for thrombin’s ability to regulate platelet function and to stimulate mitogenesis, represents a paradigm shift in understanding the general mechanism whereby proteinases regulate tissue function (Rasmussen et al., 1991; Vu et al., 1991; Hollenberg and Compton, 2002; Coughlin, 2005; Adams et al., 2011; Alexander et al., 2013, p. 1552; Hollenberg et al., 2014). A key finding was the discovery of the tethered ligand (TL) mechanism of activation of these proteinase-activated receptors (PARs) (Vu et al., 1991; Chen et al., 1994). Thus, cleavage of the N-terminus of PAR1 by thrombin at a target arginine/serine bond unmasks a TL sequence (SFLLRN—), which becomes the receptor-activating ligand (Vu et al., 1991). Similarly, thrombin cleaves an arginine-glycine bond in human PAR4 to reveal the receptor-activating TL sequence (GYPGQV—). The higher potency with which thrombin activates PAR1 versus PAR4 is due to a hirudinlike sequence in PAR1 that increases the affinity of thrombin for PAR1 relative to PAR4. Trypsin, known to target PAR2 at nanomolar concentrations (Nystedt et al., 1994), can also activate PAR4 with a potency equivalent to that of thrombin. However, in the human platelet, PAR3 acts as a cofactor for PAR4 to increase thrombin’s potency for PAR4 activation (Ishihara et al., 1997; Nakanishi-Matsui et al., 2000). Thus, both trypsin and thrombin can regulate PAR4, whereas PAR2 is preferentially activated by trypsin but not thrombin, presumably because of a lack of a thrombin-preferred upstream
proline in PAR2, just before its arginine cleavage-activation site, and the lack of a hirudin-like thrombin binding domain found in PAR1. The prolinc residue in PAR1 and PAR4 confers thrombin sensitivity to cleave the arginine and unmask the receptor-tethered ligands. Notwithstanding, human PAR2 does indeed possess potential serine proteinase cleavage sites in its N-terminal amino acid sequence at lysine residues both upstream of and downstream from its arginine-36/serine-37 TL-generating site. Thus, in principle, thrombin might be able to cleave PAR2 at a basic residue to generate signaling. Indeed, thrombin activation of PAR2 has been reported, but this result has been attributed not to a direct activation of PAR2 by thrombin but rather to an indirect transactivation of PAR2 by the PAR1 TL unmasked by thrombin (O’Brien et al., 2000). For transactivation, the unmasked TL of PAR1 (SFLRR—) is thought to reach over and function as an activating ligand for PAR2 (Chen et al., 1994; O’Brien et al., 2000). This mechanism presumably occurs in the setting of a PAR1/PAR2 heterodimer.

For the study we report here, we hypothesized that in addition to this TL crossover mechanism involving PAR1, thrombin (at sufficiently high concentrations) can activate PAR2 directly. To test this hypothesis, we used a number of approaches, including measuring the cleavage by thrombin of a synthetic peptide representing the cleavage-activation sequence of human PAR2 and the use of a clustered regularly interspaced short palindromic repeats (CRISPR)/Cas9-targeting approach to generate a PAR1-null (P1N) human embryonic kidney (HEK) cell line (P1N-HEK) and a P1N bladder cancer cell line. These P1N cells were used to evaluate the action of thrombin on cell-expressed PAR2 for signaling [calcium, mitogen-activated protein kinase (MAPK)/extracellular-regulated protein kinase (ERK) 1/2 activation, arrestin recruitment] in the absence of PAR1. Furthermore, we developed a new approach to evaluate PAR2 cleavage by thrombin and other proteinases by designing a PAR2 cleavage reporter assay, in which a NanoLuc luciferase (Nluc)-tagged PAR2 (S36L37G36/S37LIGKVDGTSHVTGKGVT) were prepared by solid-phase synthesis at the University of Calgary Peptide Synthesis Facility (peplab@uclgary.ca) as >95% pure products, verified by high-performance liquid chromatography (HPLC) and mass spectrometry. Stock solutions of about 1 mM were prepared in 25 mM HEPES buffer, pH 7.4.

Materials and Methods

Chemicals and Other Reagents. PAR-selective receptor-activating peptides (TFLLR-NH2 (TP), for PAR1; 2-furoyl-LIGRL-NO2 (2fL) for PAR2) and the peptide sequence representing the cleavage-activation site (shown as //) of human PAR2 (SSKGRR//S37LIGKVDGTSHVTGKGVT) were prepared by solid-phase synthesis at the University of Calgary Peptide Synthesis Facility (peplab@uclgary.ca) as >95% pure products, verified by high-performance liquid chromatography (HPLC) and mass spectrometry. Stock solutions of about 1 mM were prepared in 25 mM HEPES buffer, pH 7.4.

Our data illustrate unequivocally that in isolation, PAR2 signaling (activation of calcium and MAPK signals along with β-arrestin recruitment and vascular relaxation) can be regulated by thrombin cleavage, which we show via peptide mapping can target the N-terminal sequence of human PAR2 at its canonical R36/S37 cleavage-activation site.

Cell Culture. All cell media, serum, delta surface culture flasks, and multwell dishes were purchased from Thermo Fisher Scientific (Waltham, MA). HEK293 (ATCC CRL-1573) and Kirsten-virus-transformed normal rat kidney (KNRK) (ATCC CRL-1569) cell lines were routinely grown in Dulbecco’s minimal essential medium (DMEM) supplemented with 1 mM sodium pyruvate, 10% fetal bovine serum, and 2.5 μg/ml placimac (Invitrogen/ThermoFisher, Carlsbad, CA) on Nunclon Delta Surface Cell Culture Flasks (Siski Aldrich, St. Louis, MO) at 37°C in a 5% CO2 in a humidified incubator. The cell culture procedures done routinely for the measurement of calcium signaling used enzyme-free EDTA-containing cell dissociation medium for cell passage as described in detail previously (Ramachandran et al., 2011; Mihara et al., 2013). For studies of receptor cleavage (Nluc assay) or imaging, cells were passed with enzyme-free EDTA-containing cell dissociation medium or by mild trypsinization (0.2 ml of 0.25% trypsin-EDTA: Gibco/75 cm² T-flask; Thermo Fisher Scientific, Waltham, MA) to optimize cell homogeneity on plating. After dispersal (30 seconds) the trypsin was neutralized by the addition of 10% serum-containing growth medium and cells were then re-fed routinely.
Calcium Signaling Assay. Measurements of PAR2-stimulated calcium signaling in response to thrombin and to PAR2-selective agonists (trypsin and 2fLI) were conducted essentially as described previously (Kawabata et al., 1999; Ramachandran et al., 2011; Mihara et al., 2013) using a Fluor4-NW no-wash calcium assay kit (Thermo Fisher Scientific). P1-HEK and bladder cancer-derived cell line (HTB-9) cells, which express functional PAR2, were lifted from the culture flask with EDTA-supplemented calcium-free isotonic phosphate–buffered saline (pH 7.4), washed, and resuspended at approximately 5 × 10^5 cells/ml in Hanks’ buffered salt solution (HBSS) supplemented with 10 mM HEPES, 1.5 mM CaCl_2, and 1.5 mM MgCl_2 (HBSS + buffer) containing calcium indicator dye. Agonists were added to stirred cell suspensions in a spectrofluorometer, and the fluorescence emission signal at 530 nm (excitation wavelength, 480 nm) was measured relative to the signal generated in the same cell preparation by the calcium ionophore A23187 (% A23187).

Monitoring MAPK-ERK 1/2 Activation. Thrombin-stimulated MAPK-ERK 1/2 activation was monitored in KNRRK cell lines expressing either PAR1 or PAR2 essentially as described previously using western blot procedures that visualize activated phospho-MAPK (Ramachandran et al., 2009). The PAR2-expressing KNRRK cells were grown to confluence in a 24-well plate (Thermo Scientific; 2 cm^2 area) and serum starved by incubation overnight in serum-free DMEM. Fresh serum-free medium was added the following day, and the cells were incubated for another 4 hours. Thrombin (1–20 U/ml) was diluted in serum-free DMEM and then applied to the cells at room temperature for 10 minutes, at which time the cells were rinsed with ice-cold buffer and processed for western blot detection of phospho-MAPK/ERK 1/2. Activation was quantified by densitometry (ImageJ software; National Institutes of Health, Bethesda, MD) and expressed as a ratio of phospho-MAPK/ERK 1/2 relative to the signal from non-stimulated cells (fold increase over baseline). Our past work has verified that the vector-transfected or nontransfected KNRRK cells are unresponsive for PAR1/PAR2 activation of MAPK-ERK1/2.

Cloning and Transfection. The plasmid encoding the human PAR2 receptor (wt-hPAR2) cloned in pCDNA3.1 was obtained from the DNA Sequencing Facility. The functionality of wild-type and mutated constructs were confirmed by direct sequencing (University of Calgary Human Library, and we synthesized both strands of oligonucleotides for each target sequence (F2R, HGLiha, 15861_F: caacgGATAGACATAACAAGACCCG, R: aaacCGGTCTGTTATGTGTCTATCC; F2R, HGLiha, 15862_F: caacgCTCAATTGAACACCTGCTGCA, R: aacATCGAAGGGTGTTATTGAGC; F2R, HGLiha, 15861_F: caacgACCGCGGACAATGGGGGCCCCG, R: aacCGGCCCCATGTCCCAGGCGGC) were also provided by Dr. Andries Zijlstra, Vanderbilt University. The knockout design and procedures used to derive the P1N cells from the wild-type HEK and HTB-9 cells were as described by Sanjana et al. (2014) and Shalem et al. (2014) using the GeKO CRISPR protocol (https://www.addgene.org/crispr-libraries/geekv2/). Three sets of genome-specific sgRNA sequences of PAR1 (F2R) were chosen from the GeKOv2 Human Library, and we synthesized both strands of oligonucleotides for each target sequence (F2R, HGLiha, 15861_F: caacgGATAGACATAACAAGACCCG, R: aaacCGGTCTGTTATGTGTCTATCC; F2R, HGLiha, 15862_F: caacgCTCAATTGAACACCTGCTGCA, R: aacATCGAAGGGTGTTATTGAGC; F2R, HGLiha, 15861_F: caacgACCGCGGACAATGGGGGCCCCG, R: aacCGGCCCCATGTCCCAGGCGGC). Here, the upper case sequences are target sequences of the PAR1 genomic locus and the lower case sequences are flanking sequences for cloning. Both strands of the oligonucleotides were annealed and inserted in BamBI restriction enzyme sites under a U6 promoter in a CRISPR/CAS9 vector, lentiCRISPR v2, a gift from Feng Zhang (Addgene plasmid # 52961; Cambridge, MA).

The lentiCRISPR v2 plasmids containing three PAR1-targeting sequences were mixed and transfected using Lipofectamine LTX Reagent (Thermo Fisher Scientific, Waltham, MA). The transfected cells were maintained in the presence of 5 μg/ml puromycin to select knockout cells. The absence of functional PAR1 in the P1N Lenti-X 293T cells (P1N-HEK) and the P1N-HTB-9 cells was verified using a calcium signaling assay (Kawabata et al., 1999; Mihara et al., 2013), in which no signal was observed for either 1 U/ml thrombin or 25 μM of the PAR1-activating peptide, TF, in cells that were otherwise responsive to the PAR2-activating peptide, 2fLI (Fig. 3B). Furthermore, the PAR1 cells derived in this way became PAR1 responsive when reconstituted with wild-type PAR1. The Lenti-X 293T clone (P1N-HEK) and a clone of P1N-HTB-9 cells were then used as background cells to interrogate the impact of thrombin action on PAR2, which was expressed either constitutively in the cells (HTB-9) or supplemented (HEK cells) by transiently transfecting into the cells’ wild-type PAR2 either with a C-terminal eYFP (Ramachandran et al., 2009) or as a dual-tagged mRFP and C-terminal YFP: RFP-PAR2-YFP. The PAR2-expressing P1N cells (P1N-HEK/PAR2-Y; P1N-HTB9) were then used to evaluate the ability of thrombin to regulate PAR2.

PAR Cleaving Activity Monitored by an Nluc Release Assay. We developed a new assay to detect the proteolytic cleavage of the N-terminus of PAR2 by enzymes that can target either the cleavage-activation site or other extracellular locations. Our approach involves placing an N-terminal Nluc tag (Hall et al., 2012) on PAR2 that also has a C-terminal eYFP tag to monitor receptor expression (Nluc-PAR2-eYFP). Thus, our method can assess the PAR-cleaving activity.
activity of either a specific protease or any proteolytic enzyme present in body fluids or secreted by cells into a culture medium. For this reason, we sought an indicator cell in which the Nluc-tagged PAR2 could be expressed in the absence of cell-secreted proteases. To this end, we selected a human embryonic lung fibroblast cell line transformed with SV40, WI-38 VA-13 2RA (WI38VA13), as a transfected Nluc-PAR2-eYFP-expressing indicator cell, which is similar to a normal fibroblast phenotype and was found to secrete minimal PAR-protease cleavages. The Nluc-hPAR2-eYFP plasmid DNA was transfected into the WI-38 VA-13 cells with FuGENE HD (Promega Corporation) and selected with 300 μg/ml G418. The Nluc-PAR1-eYFP-expressing cells, WI38 (Nluc-PAR2), were further selected by flow cytometry (Flow Cytometry Core Facility, Health Sciences Centre, University of Calgary) by monitoring fluorescence from the C-terminal PAR2 eYFP.

The Nluc release assay to detect PAR-proteolytic activity was performed using 24-well (1-cm-diameter) plates as follows. Confluent WI38 (NlucPAR2) cells in a 775 flask were first washed three times with 1 ml of isotonic Dulbecco’s calcium/magnesium-free phosphate—buffered saline, pH 7.4, 1 mM EDTA. Ten minutes after the wash and incubation of the monolayer at 37°C in the residual buffer adhering to the monolayer, approximately 200–240 μl (4 drops with a polyethylene transfer pipette) of 0.35% (w/v) trypsin in phosphate—buffered saline, 1 mM EDTA was added to the washed monolayer and allowed to incubate for 5 minutes at room temperature to ensure disaggregation of the cells (the final trypsin concentration was approximately 0.025% w/v). Cells were then suspended in 18 ml of culture medium (DMEM/10% fetal bovine serum) to neutralize the trypsin. Aliquots of the cell suspension (500 μl) were then added into each well of a 24-well (1-cm-diameter) plate and cultured for approximately 48 hours. The cell monolayers were then washed three times with HBSS, pH 7.4, and 100 μl of 10 mM HEPES, pH 7.4-supplemented HBSS containing 1.5 mM each of MgCl2 and CaCl2 (HBSS+) was added to each well. The test amount of protease in a final volume of 100 μl of HBSS+ was then added to the Nluc-expressing cell monolayers (final volume, 200 μl) and incubated for 30 minutes at room temperature, at which time aliquots of the supernatant (100 μl) were withdrawn to measure released Nluc activity.

Aliquots from three replicate samples for each enzyme concentration were transferred to a white 96-multiwell black plate followed by the addition of diluted luciferase assay substrate solution. The substrate solution supplied with the Nano-Glo Luciferase Assay kit (Promega Corporation, Madison, WI) was diluted 10-fold with HBSS+ buffer, and 20 μl of this diluted substrate solution was added to 100 μl of cell supernatant. Luminescence from the luciferase activity, which stabilized between 5 and 60 minutes, was measured using a Victor×4 plate reader (PerkinElmer, Waltham, MA), according to the manufacturer’s instructions. The luminescence yield was approximately linear between approximately 0.5 and 10 nM benzoyl-L-arginine ethyl ester U/ml trypsin activity. For concentration-effect curves measuring the ability of thrombin and trypsin to cleave and release the N-terminal Nluc reporter, data were normalized for each enzyme concentration relative to the background luminescence observed in the absence of enzyme (% luminescence). Work was done with both wild-type Nluc-tagged PAR2 and with Nluc-tagged PAR2 having an R36G mutation that prevents cleavage and unmasking of the PAR2 TL sequence by trypsin or other serine proteases. On average, the maximum increase in luminescence over background was from 4500 to 27,400 and 2100 to 2300 luminescence counts/0.1 second/sample for wild-type and mutant PAR2 by thrombin cleavage, compared with 1400–31,000 and 500–2900 luminescence counts/0.1 second/sample for wild-type and mutant PAR2 by trypsin.

**Imaging Cleavage and Activation of Dual-Labeled PAR2.** Cleavage of the N-terminal domain of PAR2 expressed in a HEK cell background, cultured for imaging experiments as described previously, was monitored by confocal imaging using an N-terminal mRFP and C-terminal eYFP dually tagged PAR2 (RFP-PAR2Y) either with or without an alanine for arginine substitution at residue 36 (R36A: RFP-PAR2Y/R/A) to eliminate the TL site cleaved by thrombin. Monolayers that had grown for 24 hours in the matrix metalloprotease inhibitor–containing medium were washed three times with DMEM, and PAR2 activation was triggered by the addition of either enzyme (trypsin, 2–5 U/ml; thrombin, 25–50 U/ml) or PAR2-activating peptide (2 μM fligrlo-NH2) to the cell supernatant. Receptor activation was stopped at timed intervals after the addition of 1 ml of 10.5% formalin fixative (BDH Chemicals/VWR, Edmonton, Alberta, Canada) to the 0.2 ml of cell supernatant. Upon confocal imaging, intact PAR2 appears yellow, combining the red from the N-terminal mRFP and artificial green from the C-terminal eYFP. After cleavage and activation by thrombin, the N-terminal mRFP is released and PAR2 appears green due to the C-terminal eYFP tag.

**Measurement of Bioluminescence Resonance Energy Transfer between PAR2-eYFP and Arrestins 1 and 2.** Measurements of bioluminescence resonance energy transfer between C-terminal eYFP-tagged PAR2 and Renilla reniformis luciferase–tagged β-arrestins 1 and 2 were measured as described previously (Ramachandran et al., 2009, 2011; Mihara et al., 2013), with luminescence values measured 20 minutes after receptor activation, in keeping with the procedure described by Hamdan et al. (2005).

**Measurements of PAR-Induced Endothelium-Dependent Vascular Relaxation.** Endothelium-dependent vasorelaxation was measured as described previously (McGuire et al., 2002) using isolated mouse aorta rings obtained from P11 mice used in previous studies of PAR function (Cenac et al., 2005). Tissues with a verified endothelium-dependent relaxant response to acetylcholine were constricted with phenylephrine (2.5 μM), and a relaxant response to thrombin (5–50 U/ml) or a PAR2-activating peptide (2fLI) was monitored. The response was measured in the absence and presence of a 10-fold molar excess thrombin-inhibitory concentration of hirudin. The thrombin response was also measured (10 U/ml) after first desensitizing the vascular endothelial response by repeated exposure to the PAR2-activating peptide, 2fLI (2 × 5 μM).

**Results and Discussion**

**Thrombin Cleavage of a Synthetic Peptide Derived from the N-terminal PAR2 TL Sequence.** We first established that thrombin can cleave a synthetic peptide derived from the N-terminal TL-containing peptide sequence, SSKGRGLIGRLO-NH2 (TL sequence unmasked by cleavage (//) shown in red). The synthetic peptide (100 μM) representing the N-terminal cleavage-activation domain of PAR2 was subjected to thrombin cleavage (25 U/ml) and HPLC analysis followed by matrix-assisted laser desorption/ionization mass spectral identification of the peptide fragments as described in Materials and Methods.
from the N-terminal human PAR2 that contains the canonical PAR2 TL sequence unmasked by trypsin (GTNRSSKG/SLIKVGDTSHTGKVGTV, where the trypsin cleavage site denoted by //; TL sequence in bold). As illustrated in Fig. 1, thrombin (25 U/ml) can indeed cleave the synthetic PAR2-derived N-terminal peptide to unmask the receptor-activating TL sequence (SLIKV–). Of note, although not identified as a distinct peak in the HPLC analysis, deconvolution of the peptide masses detected by matrix-assisted laser desorption/ionization mass spectral analysis of the cleavage reaction mixture provided evidence for the presence of the peptide SSKGRSSLIKVGDTSHTGKVGTV. That peptide would originate from an upstream cleavage of the PAR2 sequence: GTNR//SSKGRSSLIKVGDTSHTGKVGTV. The next issue to deal with was to verify that thrombin is able to cleave the N-terminal sequence of PAR2 when expressed in an intact cell system.

**Thrombin Cleavage and Release of Luciferase Activity from the Nluc-Tagged-PAR2 N-Terminus.** We monitored the ability of thrombin to cleave the PAR2 N-terminus, as indicated by the release of the N-terminal Nluc tag from PAR2 expressed in WI38VA13 cells (shown in Fig. 2A). As shown in Fig. 2, B and C, thrombin, like trypsin (2–50 U/ml of each enzyme), was able to release the N-terminus of PAR2 from the cell surface to yield a luciferase signal in the supernatant. Thrombin was also able to cleave and release the PAR2 N-terminal domain when the receptor was expressed in several other cell backgrounds (e.g., Chinese hamster ovary; data not shown). Significantly, the Nluc-PAR2 clone with an R36G mutation at the trypsin cleavage-activation site, which is no longer a tryptic substrate, did not release luciferase activity when treated with either trypsin or thrombin at concentrations up to 2 U/ml trypsin and 50 U/ml thrombin activity (dashed lines in Fig. 2, B and C). Given that thrombin appeared to cleave the N-terminus of PAR2 at the same arginine-36 as trypsin, the next issue was to determine if the cleavage resulted in receptor activation.

**Activation of PAR2 by Thrombin to Stimulate Calcium and MAPK Signaling along with β-Arrestin Interactions.** Since previous publications indicate that PAR1 and PAR2 can cooperate for cell signaling, possibly as a heterodimer, we wished to use cells in which PAR2 can signal in the absence of PAR1. Therefore, as done in previous work (Al-Ani et al., 2004), we used PAR2-transfected KNRK cells that as a wild-type cell do not express either functional PAR1 or PAR2 and that as transfected cells express functional PARs. Furthermore, we used a HEK293 background cell line that constitutively expresses functional PAR1 and PAR2 (Kawabata et al., 1999) to prepare a P1N-HEK cell line with a CRISPR/Cas9 approach. The P1N-HEK cells were also used as transient transfection targets for the expression of either PAR2 with a C-terminal eYFP tag or dually tagged PAR2, having an N-terminal RFP tag and a C-terminal eYFP. In the P1N-HEK cell background, these tagged PAR2 constructs were used to monitor the following: 1) PAR2-mediated calcium signaling activated by thrombin, trypsin, or a PAR2-activating peptide; 2) interactions of activated PAR2 with β-arrestins 1 and 2, as described previously (Ramachandran et al., 2011); and 3) receptor cleavage and postactivation dynamics, as done previously for PAR1 (Mihara et al., 2013). The PAR2-expressing KNRK cells were used to monitor PAR2-mediated MAPK signaling triggered by thrombin, in keeping with work done in the past using these PAR2-expressing cells (Ramachandran et al., 1999).

**Calcium Signaling.** As shown in the left panel in Fig. 3A, the wild-type Lenti-X 293T cells (wild-type HEK), from which the P1N-HEK line was derived, generated a prominent calcium signal when stimulated by either the PAR1-selective agonist, TF, or the PAR2-selective agonist, 2fLI. Furthermore, a calcium signal was stimulated by either thrombin (via PAR1) or trypsin (via PAR2) in the wild-type HEK 293T cells that had first been cross-desensitized by pretreatment with the selective PAR1 agonist (TF) to enable activation of PAR2 only, or desensitization with the selective PAR2 agonist (2fLI) to measure only PAR1 activation. Thus, thrombin yielded a PAR1 calcium signal in the PAR2-desensitized cells; and trypsin likewise yielded a PAR2 signal in the PAR1-desensitized cells. In contrast, at concentrations that activated PAR1 in the wild-type HEK293 cells, neither the PAR1-activating peptide, TF (10–25 μM), nor thrombin (25–50 U/ml) was able to generate a calcium signal in the P1N-HEK cells, whereas the cells were fully responsive to PAR2

![Fig. 2. Thrombin cleavage and release of the PAR2 N-terminal domain from Nluc-tagged PAR2.](image)

(A, left) The scheme illustrates the mechanism whereby thrombin cleavage of the N-terminus of PAR2 releases Nluc into the cell supernatant as an index of receptor cleavage. (B and C, middle and right) Concentration-effect curves for the cleavage and release of the N-terminal domain of PAR2 by increasing concentrations of either thrombin (National Institute of Health U/ml) (B) or trypsin (Nα-Benzoyl-L-arginine ethyl ester U/ml) (C). Cleavage of the wild-type PAR2 sequence (solid line) is compared with cleavage of the R36G mutant (dashed line), which is resistant to serine protease cleavage. Values (% luminescence, relative to control untreated monolayer values) represent the mean values ± S.E.M. (bars) for three replicates within an individual Nluc cleavage assay. Baseline absolute luminescence values ranged from 1400 (trypsin) to 4500 (thrombin) luminescence units for wild-type PAR2 and 455 (trypsin) to 2000 (thrombin) luminescence units for the RA mutant PAR2 and maximal values were from 36,000 (trypsin, 50 U/ml) to 27,000 units (thrombin, 50 U/ml) for wild-type PAR2.
activation either by the PAR2-activating peptide, 2fLI, or by trypsin (left-hand panel in Fig. 3B). However, when the P1N-HEK cells were transiently transfected with dual-tagged PAR2 to increase the abundance of PAR2 expression, thrombin was able to stimulate a calcium signal that was about 40% of the magnitude of the signal triggered by trypsin (left-hand panel in Fig. 3C). In the same P1N-HEK cells in which PAR2 was desensitized, the PAR1-activating peptide, 2fLI (Thr, 25–50 U/ml), but not to the PAR1-activating peptide (Thr, 10–25 μM TF) added after the cells were first desensitized with 2fLI to eliminate TF-mediated PAR2 signaling. (D) P1N-HEK cells transfected with the R36A mutant of YFP-labeled PAR2 (P1N-HEK/PAR2-2fLI-Y) respond to 2fLI (Thr, 1 μM) but do not respond to thrombin (Thr, 25–50 U/ml) or to the PAR1-activating peptide TF (Thr, 25 μM TF; added to cells pretreated with 2fLI to eliminate the TF-mediated PAR2 calcium signal). (E) Hirudin (Thr, 2.5 μM) blocks the PAR2 response to thrombin (Thr, 25 U/ml). (F) Desensitizing P1N-HEK/PAR2-Y-expressing cells by 2× treatment with 2fLI (Thr, 5 μM) desensitizes the PAR2 response to thrombin (Thr, 25 U/ml). (G and H) Right-most panels: (G) Concentration-effect curves for thrombin activation of PAR2 calcium signaling in P1N-HEK cells expressing PAR2 (P1N-HEK/PAR2-Y), however, the PAR2-activating peptide, which acts independently of the tethered ligand, was still able to trigger a calcium signal (second response from the left in the left-hand panel in Fig. 3D). This result demonstrated the presence of ample functional R36A-PAR2 in the cells that were resistant to enzyme cleavage and activation but not to activation by the PAR2-activating peptide (Fig. 3D). Thus, thrombin yielded a PAR2 signal in the P1N-HEK cells only when transfected with a PAR2 construct with a susceptible tryptic cleavage site at arginine-36.

To validate the data obtained with the P1N-HEK cells, we generated a P1N-HTB-9 cell line. In the wild-type HTB-9 cells, both PAR1 and PAR2 agonists caused calcium signals equivalent to those observed for the wild-type HEK cells
was abrogated both by predesensitization of the PAR2 preparation from the P1N animals due to PAR2 activation in P1N mice (Fig. 3H). The relaxant response of the aorta PAR2-responsive aorta tissue preparation derived from P1N-HEK and P1N-HTB-9 cells that express functional PAR2. Thus, thrombin at sufficiently high concentrations can, in principle, signal via both PAR1 and PAR2 using both a direct cleavage mechanism as well as an indirect (i.e., TL crossover) mechanism for PAR2 activation.

**Vascular Action of Thrombin via PAR2 in P1N Mice.** To evaluate the potential impact of thrombin to activate endothelial PAR2 in vivo, we measured the endothelium-dependent relaxant activity of thrombin in a PAR2-responsive aorta tissue preparation derived from the P1N animals due to PAR2 activation was abrogated both by predesensitization of the PAR2 relaxant response with repeated exposure to the PAR2 agonist 2fLI (second tracing on the right in Fig. 3H) and by neutralizing the thrombin preparation with a 10-fold molar excess of hirudin (data not shown). Furthermore, the addition of 1 μg/ml soya trypsin inhibitor did not affect the thrombin-mediated relaxant response but completely blocked the response to 0.5 U/ml trypsin (data not shown). The results with the enzyme-selective inhibitors of thrombin and trypsin verified that the relaxation was generated by thrombin and not by a contaminating trypsin-like enzyme. Our data thus showed that thrombin is capable of causing PAR2 activation in intact vascular tissues in vivo, albeit at relatively high concentrations relative to those that cause coagulation.

**MAPK-ERK 1/2 Activation.** In addition to stimulating PAR2 calcium signaling in the P1N-HEK cells, thrombin also stimulated MAPK-ERK 1/2 signaling in the PAR2-expressing KNRK cells (Fig. 4). The phospho-MAPK signal caused by 2–20 U/ml thrombin in the PAR2-expressing KNRK cells (left panel in Fig. 4A) was comparable to the signal generated by 1 U/ml thrombin in the P1N-expressing KNRK cells (right panel in Fig. 4A). The densitometrically quantified concentration dependence of thrombin-mediated PAR2 activation is shown in Fig. 4B.

**Thrombin-Stimulated PAR2–Arrestin Interactions.** Using the P1N-HEK cells expressing N-RFP/C-YFP-tagged PAR2, we next evaluated the ability of thrombin to drive PAR2–arrestin interactions. As shown in Fig. 5, thrombin-mediated activation of PAR2 promoted interaction of the receptor with both arrestins 1 and 2. We thus concluded that at concentrations in the range of 10–25 U/ml (100–250 nM), thrombin can stimulate PAR2 calcium and MAPK signaling along with β-arrestin interactions.

**Visualizing Thrombin-Triggered PAR2 Cleavage and Activation.** Finally, we wished to visualize PAR2 cleavage and activation by thrombin using confocal microscopy. As shown in panel 2 in Fig. 6, thrombin cleavage released the

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**Fig. 4.** Thrombin activation of PAR2 stimulates MAPK signaling in PAR2-expressing KNRK cells (KNRK-PAR2). Using previously described methods (Ramachandran et al., 2009; Mihara et al., 2013), KNRK cells, which do not respond to thrombin, were transfected with either PAR2 (left-hand western blot) or PAR1 (right-hand western blot), and the PAR-expressing cell lines so derived were treated with thrombin (2 or 20 U/ml). (A) Western blots: cells were lysed after 10 minutes, and the activation of MAPK was measured by western blot analysis (P-MAPK) as described previously. (B) Densitometry of bands: P42/44 phospho-MAPK-ERK 1/2 (p-ERK 1/2) bands were quantified densitometrically, and MAPK-ERK 1/2 activation was expressed as an increase relative to the signal observed in untreated samples (factor increase in p-ERK 1/2 over baseline). Histograms represent the average fold increase ± S.E.M. (bars: P ≤ 0.05 for all histogram values compared with untreated samples) for three replicate experiments.

**Fig. 5.** Thrombin-mediated PAR2–β-arrestin interactions. PAR2, C-terminally tagged with eYFP (P2Y) was transfected into the P1N-HEK cells (P1N-HEK/P2Y) along with Renilla reniformis luciferase–tagged arrestin 1 or 2. Bioluminescence resonance energy transfer between the receptor and either β-arrestin 1 (Arr1) or β-arrestin 2 (Arr2) [bioluminescence resonance energy transfer (BRET) ratio NanoLuc luciferase (nLuc)/YFP] was then measured as described previously (Hamdan et al., 2005; Ramachandran et al., 2009) and in Materials and Methods. Histograms represent the average fold increase ± S.E.M. (bars: P ≤ 0.05 for all histogram values compared with thrombin-untreated samples) for three replicate measurements on independently grown cell samples.
background HEK and HTB-9 cells in vitro (0.5–5 U/ml; 5–50 nM) (Mihara et al., 2013).

Although our data fully support our hypothesis that thrombin at sufficiently high concentrations can activate PAR2 directly, the following question remains: Can this action of thrombin play a physiologic role? We suggest that the answer to this question is yes, but only in unique situations such as in a tumor microenvironment or in the setting of acute tissue trauma, where high levels of active thrombin can be generated. Thus, since the circulating concentration of prothrombin in normal subjects ranges from 700 nM to 1.7 μM (Henderson et al., 1980), its conversion to thrombin in an acute setting could indeed reach the levels we find can activate PAR2. In fact, unusually high levels of acutely generated thrombin have been found in vivo in cancer patients. In a study of 1033 patients with malignancies of the breast, lung, gastrointestinal tract, pancreas, kidney, prostate, or brain—or having lymphoma, multiple myeloma, or other tumor types—the median peak thrombin concentration was found to be 500 nM (Ay et al., 2011). Similarly, breast cancer patients have demonstrated significantly higher levels of thrombin generation than controls, reaching levels of greater than 500 nM (Chaari et al., 2014). Moreover, in a study of blunt trauma patients, the range of peak thrombin in the injured individuals was nearly 30 times that of controls, and the patients with the most severe injury had thrombin levels of 300 nM (Park et al., 2012). Thus, the thrombin concentrations we have shown that can activate PAR2 can indeed be found in vivo.

In addition, although PAR1 and PAR2 are often coexpressed in a cellular setting, there are situations in which PAR2 may be expressed in the absence of PAR1. In that circumstance, the ability of an activated thrombin receptor to transactivate PAR2 would not be possible; therefore, a direct activation of PAR2 by thrombin would be of physiologic significance. Thus, our data indicate that one can add PAR2 to the list of thrombin targets that can potentially regulate cell signaling in addition to PAR1 and PAR4. The impact of this action of thrombin, which we propose is feasible in a restricted environment where a substantial fraction of circulating prothrombin is activated (e.g., in a tumor microenvironment), remains to be established in vivo.

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Authorship Contributions

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