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**Neuronal FcγRI mediates acute and chronic joint pain**

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Abstract

Although joint pain in rheumatoid arthritis (RA) is conventionally thought to result from inflammation, arthritis pain and joint inflammation are at least partially uncoupled. This suggests that additional pain mechanisms in RA remain to be explored. Here we show that FcγRI, an immune receptor for IgG immune complex (IgG-IC), is expressed in a subpopulation of joint sensory neurons and that, under naïve conditions, FcγRI crosslinking by IgG-IC directly activates the somata and peripheral terminals of these neurons to evoke acute joint hypernociception without obvious concurrent joint inflammation. These effects were diminished in both global and sensory neuron-specific Fcgr1 knockout mice. In murine models of inflammatory arthritis, FcγRI signaling was upregulated in joint sensory neurons. Acute blockade or global genetic deletion of Fcgr1 significantly attenuated arthritis pain and hyperactivity of joint sensory neurons without measurably altering joint inflammation. Conditional deletion of Fcgr1 in sensory neurons produced similar analgesic effects in these models. We therefore suggest that FcγRI expressed in sensory neurons contributes to arthritis pain independently of its functions in inflammatory cells. These findings expand our understanding of the immunosensory capabilities of sensory neurons and imply that neuronal FcγRI merits consideration as a target for treating RA pain.

Keywords: pain; FcγRI; hyperalgesia; rheumatoid arthritis, inflammation, dorsal root ganglion.
Introduction

Rheumatoid arthritis (RA) is a chronic autoimmune disorder that affects millions of American adults and exhibits high morbidity and mortality. Joint pain is a cardinal clinical feature of RA, and poses an enormous health burden (1, 2). RA is characterized by synovitis and joint destruction, both of which are significant contributors to the associated joint pain. Accordingly, most current treatments for RA pain are aimed at reducing joint inflammation and slowing joint damage. Yet, many of these anti-inflammatory therapies exhibit limited efficacy and/or adverse side effects (3, 4). While RA pain is often viewed simply as a direct consequence of inflammation, pain and inflammation are at least partially uncoupled in RA (5, 6). Joint pain often precedes overt signs of joint inflammation and can persist in subpopulations of RA patients despite seemingly optimal control of inflammation with current biologic therapies (7-9). A recent study demonstrated that anti-citrullinated protein antibodies (ACPA) may trigger RA pain via an inflammatory cell-independent mechanism that involves release of the nociceptive chemokine CXCL1/IL-8 from osteoclasts (10). Thus, additional mechanisms besides inflammation likely contribute to RA pain. Yet, such mechanisms remain largely unexplored.

One important pathological entity in RA is immunoglobulin G-immune complex (IgG-IC), which is present at high amounts in the serum and affected joints of RA patients (11, 12). IgG-IC exerts its biological effects largely through Fc gamma receptors (FcyRs), which are most prominently expressed in immune cells and are critical regulators of immunity (13-16). Of four FcyR subtypes (I-IV) in rodents, most are low-affinity receptors and only bind IgG in ICs. However, FcyRI (also called CD64) is the sole high-affinity receptor that can bind both monomeric and polymeric IgG (16). IgG-IC/FcyRI signaling has been implicated in the pathogenesis of RA. Fcgr1 deficient mice exhibited decreased arthritic symptoms in collagen-induced arthritis (CIA) and antigen-induced arthritis (AIA) models (11, 17, 18) and treatment with a CD64-directed immunotoxin diminished inflammation and bone erosion in human CD64 transgenic rats suffering from joint inflammation (19). Similarly, scavenging IgG-IC with
recombinant soluble FcγRI suppressed cartilage destruction in AIA and CIA models (11, 20). However, FcγRI plays differential roles in arthritis pathogenesis in different animal models of RA. Whereas FcγRI is of crucial importance in both severe joint inflammation and cartilage destruction in the CIA model (11, 20), in the AIA model this receptor predominantly mediates cartilage destruction without a clear role in joint inflammation (17, 18). Thus, the AIA model enables us to focus on the potential roles of FcγRI in AIA-associated pain beyond those in inflammation, per se.

Although IgG-IC/FcγRI signaling has been suggested to play a prominent role in arthritis pathogenesis (11, 17, 18, 21), there are no reports specifically addressing its potential contributions to RA pain. Given that FcγRI is widely expressed in immune cells (13-16), FcγRI has been thought to contribute to RA pain by inducing proinflammatory cytokine release from immune cells. While multiple cytokines appear to sensitize joint nociceptors (22-26), this cannot explain all components of RA pain. Our group and others have revealed that, in addition to its expression in immune cells, FcγRI, but not subtypes II or III, is also expressed in subsets of nociceptive dorsal root ganglion (DRG) neurons of rats and mice (27, 28). These neuronal receptors are functional, since in DRG neuron culture, IgG-IC directly induces neuronal activation through a process that involves FcγRI and a downstream ion channel, TRPC3 (29). Yet, the in vivo relevance of these findings has not been explored, and no studies have addressed whether FcγRI is expressed in joint sensory neurons or whether IgG-IC acts directly on joint sensory afferents through neuronal FcγRI to induce joint pain. It is also unknown whether neuronal FcγRI contributes to joint hypernociception in the setting of arthritis. In this study, we tested the hypothesis that IgG-IC signaling contributes to RA joint pain, at least in part, through a mechanism involving direct activation of neuronal FcγRI.

Results

FcγRI is expressed in mouse joint innervating DRG neurons
Since all commercially available anti-FcγRI antibodies that we tested lacked adequate specificity for reliable immunostaining of mouse tissues, we alternatively performed in situ hybridization (ISH) to map the expression pattern of FcγRI in joint innervating DRG neurons that were retrogradely labeled by fast blue (FB) injection into the ankle joint. ISH revealed Fcgr1 mRNA expression in 27.5% of FB-labeled joint sensory neurons, regardless of cell size (Figure 1A). Fcgr1 ISH signal was colocalized with immunostaining for the neuronal-specific nuclear protein NeuN (Figure 1A), suggesting neuronal expression of Fcgr1 mRNA. The specificity of Fcgr1 mRNA detection was validated by a loss of ISH signal in the DRG of global Fcgr1−/− mice (Figure 1B) and in wildtype (WT) DRG sections stained with a sense control probe (Figure 1C).

To further define the expression pattern of FcγRI in joint sensory DRG neurons, we performed double staining for FcγRI and markers of different cell populations. Among FB-labeled joint sensory neurons, Fcgr1 mRNA expression was detected in both small- (peripherin+) and large-diameter (NF200+) neurons (Figure 1D). In addition, 53.3% of Fcgr1+ neurons co-expressed CGRP, a marker for nociceptive peptidergic neurons (Figure 1D). However, we did not detect obvious colocalization of Fcgr1 mRNA expression with glutamine synthetase (GS), a satellite glial cell marker (Figure 1D). These results indicate that a subpopulation of joint sensory neurons, including nociceptors, express FcγRI, providing an anatomical basis for neuronal FcγRI modulation of joint pain.

**IgG-IC directly activates joint sensory afferents through neuronal FcγRI**

To examine whether the FcγRI expressed in joint sensory neurons is functional, we investigated the effects of IgG-IC on Ca2+ responses in dissociated Dil-labeled DRG neurons from WT(Fcgr1+/+) and global Fcgr1−/− mice using ratiometric Ca2+ imaging. In Fcgr1−/− mice, application of IgG-IC (1 µg/ml), but not antigen (BSA) or antibody alone (anti-BSA IgG), evoked Ca2+ increases in 17.6% of joint sensory neurons (Figure 2A, B). Moreover, 80% of IgG-IC responsive neurons responded to capsaicin. In contrast, IgG-IC evoked Ca2+ Responses in a
significantly smaller fraction (3.8%) of joint sensory neurons from global Fcγr1−/− mice (Figure 2A, B).

To visualize whether IgG-IC directly activates peripheral terminals of joint sensory afferents through FcγRI, we performed in vivo imaging on the cell bodies of retrogradely Dil-labeled DRG neurons of mice (PirtCre-GCamp6) which express the fluorescent calcium indicator GCAMP6 in peripheral sensory neurons (30). Injection of IgG-IC (100 µg/ml; 10 µl), but not vehicle (PBS) or monomeric IgG, to the right hind ankle joint cavity evoked Ca²⁺ responses in a subset of Dil-labeling joint sensory neurons (Figure 2C,D). Moreover, all IgG-IC responsive neurons responded to mechanical stimulation when the ankle was pressed with blunt forceps (Figure 2C). However, the proportion of IgG-IC-responsive neurons was significantly diminished in PirtCre-GCamp6 mice crossed onto global Fcγr1−/− mice (Figure 2C,D). These findings support the idea that IgG-IC has a direct excitatory action on joint sensory afferents via the activation of neuronally expressed FcγRI.

Local administration IgG-IC elicits acute joint pain-related hypersensitivity without obvious inflammation in naïve mice

Given that IgG-IC is able to stimulate immune cells, directly activate joint sensory neurons, or both, we assessed the effects of IgG-IC at different doses on joint pain and inflammatory processes in vivo. Intraarticular (i.a) injection of IgG-IC, but not the vehicle (PBS) or monomeric IgG, into the right hind ankle cavity of the mouse significantly reduced mechanical response threshold in the hind ankle, and increased hindpaw withdrawal frequency to mechanical stimulation of nearby glabrous paw skin in a dose-dependent manner (Figure 3A-C). No secondary heat hyperalgesia in the hindpaw skin or obvious joint swelling were observed after i.a injection of IgG-IC (Figure 3D,E). The pro-nociceptive effects lasted for at least 5 h after injection and were resolved 24 h later. Since IgG-IC evoked nociceptive behaviors may result from joint inflammation, we performed qRT-PCR to assay for changes in a subset of
inflammatory markers in the synovium at the early (1 h) and late phases (5 h) following i.a.
injection of IgG-IC. However, no significant differences were observed in the mRNA expression
levels of cytokines (\textit{Tnfa}, \textit{Il6}, \textit{Il1b}), chemokines (\textit{Mcp1}, \textit{Cxcl1}), matrix metalloproteases
(\textit{Mmp2,9,13}), T cell (\textit{Cd3}) or macrophage (\textit{Cd68}) markers, or mast cell proteases (\textit{Mcpt4},
\textit{Tpsb2}) in the joint synovium between treatments at 1 h (Supplemental Figure 1A) or 5 h
(Supplemental Figure 2A) after injection. To further determine whether i.a injection of IgG-IC
induced a local immune response, we employed immunohistochemical (IHC) staining to assess
cellular infiltration of macrophages, neutrophils, lymphocytes, and mast cells in the joint
synovium, using the markers CD68, Ly6G/C, CD3, and c-Kit, respectively. No obvious
differences were observed in any of these markers between groups 1 h after the injection
(Figure 3F,G). However, the expression of Ly6G/C and CD3 in the synovium was significantly
increased at 5 h after injection in IgG-IC-treated mice, compared to those treated with vehicle or
monomeric IgG (Supplemental Figure 2B,C). Likewise, joint histological H&E staining analysis
did not reveal any signs of immune cell infiltration or synovial hyperplasia in any groups at 1h
(Figure 3H) or 5 h (Supplemental Figure 2D) after injection. In addition, toluidine blue (TB)/fast
green staining showed that IgG-IC did not cause any obvious bone or cartilage destruction in
the joint compared to controls (Supplemental Figure 1B and Supplemental Figure 2E). These
data suggest that IgG-IC is sufficient to evoke behavioral signs of acute joint pain without
concurrent inflammation, at least at early stages.

\textbf{FcγRI expressed in primary sensory neurons mediates IgG-IC-evoked nocifensive
behaviors}

Since FcγRI is a receptor for IgG-IC, we asked whether FcγRI mediates the acute
pronociceptive effects of IgG-IC using global \textit{Fcgr1}\(^{-}\) mice. Global \textit{Fcgr1}\(^{-}\) mice exhibited normal
basal mechanical sensitivity in the hind ankle and hindpaw, and normal heat sensitivity in the
hindpaw, compared to \textit{Fcgr1}\(^{++}\) littermates (Figure 4A-C). However, primary mechanical
hypersensitivity in the ankle and secondary mechanical hypersensitivity in the hindpaw upon i.a. injection of IgG-IC were significantly attenuated in Fcgr1\(^{-/}\) mice compared to Fcgr1\(^{+/}\) littermates (Figure 4D-F). These findings suggest that FcγRI is necessary for IgG-IC-elicited acute nocifensive behaviors. Given that FcγRI is widely expressed in immune cells, we next investigated whether FcγRI-bearing immune cells are required for IgG-IC-induced joint hypernociception. In naïve mice, i.a injection of clodronate-laden liposomes, but not control liposomes produced optimal depletion of synovial lining macrophages (31, 32), but did not affect basal mechanical or heat nociception 7 days after injection (Supplemental Figure 3A-E). In addition, IgG-IC-evoked nocifensive behaviors were not significantly different between liposomal clodronate-treated mice and those treated with control liposomes (Figure 4G,H). In mouse strains lacking either T cells (Rag1\(^{-/}\)) or mast cells (c-Kit\(^{Wsh/Wsh}\)) we observed no significant differences in basal mechanical or thermal sensitivity (Supplemental Figure 3F-K) or in IgG-IC-evoked nocifensive behaviors, compared to WT controls (Figure 4I-L). Together, these results indicate that the pronociceptive effects of IgG-IC are mediated by FcγRI, but do not specifically require macrophages, lymphocytes or mast cells. This left open the possibility that IgG-IC acts directly on FcγRI expressed on joint innervating sensory neurons to elicit articular hypernociception via non-immune modulation.

To more directly assess the potential involvement of neuronal FcγRI in the pronociceptive effects of IgG-IC, we generated a new mouse line bearing a conditional deletion allele of Fcgr1 (Fcgr1\(^{fl/fl}\)) and crossed these mice with the PirtCre line to selectively omit Fcgr1 expression from peripheral sensory neurons (Figure 5A). qRT-PCR and ISH analysis in PirtCre; Fcgr1\(^{fl/fl}\) mice confirmed that loss of Fcgr1 expression specifically occurred in the DRG but not the spleen (Figure 5B,C). Adult PirtCre; Fcgr1\(^{fl/fl}\) mice did not exhibit any abnormalities in basal sensitivity to mechanical or heat stimuli applied to the ankle or plantar skin of hindpaws compared to WT littermate controls (Figure 5D-F). However, PirtCre; Fcgr1\(^{fl/fl}\) mice exhibited less primary mechanical hypersensitivity in the ankle and less secondary mechanical hyperalgesia in the
hindpaw following i.a injection of IgG-IC, compared with *PirtCre* negative *Fcgr1*+/− control littermates (Figure 5G-I). These results support the notion that neuronal FcγRI contributes to IgG-IC- evoked acute nocifensive behaviors in the naïve state.

**AIA upregulates FcγRI expression and function in DRG**

We next employed the well-established AIA murine model of RA, in which the provocative antigen is methylated bovine serum albumin (mBSA) (33, 34). We performed qRT-PCR on DRG tissue from WT mice subjected to this model to assay for alterations of *Fcgr1* mRNA expression. The *Fcgr1* mRNA expression level in the DRG was significantly greater in AIA mice than vehicle control mice on days 1 and 3 after challenge (Figure 6A,B). ISH analysis further revealed that, 3 days after challenge, a larger percentage of DRG neurons (including FB-labeled joint and non-FB labeled non-joint sensory neurons) expressed *Fcgr1* mRNA in AIA mice, compared to vehicle control–treated animals (Figure 6C-E). Moreover, a greater proportion of *Fcgr1*+ joint sensory neurons in AIA mice co-expressed CGRP, as compared to those in control mice (Figure 6B,F). To determine whether the function of FcγRI in joint sensory neurons was enhanced in the context of arthritis, we compared Ca²⁺ responses evoked by IgG-IC in Dil-labeled joint innervating DRG neurons from control versus AIA mice using ratiometric Ca²⁺ imaging. As expected, the percentage of IgG-IC responsive neurons was greater in AIA mice than that in control animals (Figure 6G). These findings suggest that the expression and function of FcγRI in joint sensory neurons is significantly upregulated in the setting of AIA.

**Genetic deletion of *Fcgr1* attenuates arthritis pain in inflammatory arthritis models without obvious effects on joint inflammation**

To explore whether FcγRI contributes to arthritis pain, we compared pain-related behaviors between *Fcgr1*+/+ and *Fcgr1*−/− mice following AIA. Although both *Fcgr1*+/+ and *Fcgr1*−/− mice developed primary mechanical hyperalgesia in the hind ankle, this effect was attenuated in
global \(Fcgr1^{-/-}\) compared with \(Fcgr1^{+/+}\) mice (Figure 7A). Similarly, global \(Fcgr1^{-/-}\) mice exhibited less secondary mechanical and thermal hyperalgesia in the hindpaw than WT littermates over the course of AIA (Figure 7B-D). However, no significant differences in joint swelling following AIA were observed between genotypes (Figure 7E). To further determine whether the apparent antihyperalgesic effects of \(Fcgr1\) knockout were attributable to a possible attenuation of joint inflammation, we measured the mRNA expression of a number of inflammatory mediators in the synovium 7 h and 1 day following AIA. Among all the genes tested, AIA caused significant upregulation of \(Il1b\), \(Il6\), \(Tnfa\), and \(Cxcl1\) in the synovium 7 h after AIA (Supplemental Figure 4A). However, there were no significant differences in the alterations of the mRNA expression levels of these cytokines between genotypes. Similarly, the mRNA expression levels of \(Mcp1\) and \(Mmp9\), in addition to \(Il1b\), \(Il6\), and \(Cxcl1\), were upregulated to the same extent in both genotypes on day 1 after AIA (Supplemental Figure 4B). No significant changes in mRNA expression of other inflammatory mediators were observed in either genotype at two time points following AIA (Supplemental Figure 4B). On day 2 after AIA, joint IHC analysis showed that the marker for macrophages (CD68) was increased, but markers for T cells (CD3), neutrophils and monocytes (Ly6C/G), and mast cells (c-Kit) were not altered (Supplemental Figure 5A,B). On day 4 after AIA, all assayed immune cell markers except for c-Kit were increased (Supplemental Figure 6A-B). However, none of the AIA-induced increases in any of these markers were significantly different between genotypes (Supplemental Figure 5A,B; Supplemental Figure 6A,B). Similarly, joint histological H&E staining did not show significant differences in immune cell infiltration between genotypes on either day 2 (Supplemental Figure 5C,D) or day 4 (Supplemental Figure 6C,D) after AIA. In addition, we did not observe obvious immune cell infiltration within DRG on days 2 (data not shown) or 4 following AIA in either genotype (Supplemental Figure 7). Although FcyRI is apparently not involved in joint inflammation in the AIA model, it acts as a critical player in cartilage destruction during AIA (18, 35). Thus, we next asked whether diminished AIA-associated hypernociception in global \(Fcgr1^{-/-}\) mice is secondary to reduced cartilage
destruction. However, TB/fast green staining analysis did not detect any obvious cartilage destruction on days 2 (Supplemental Figure 8A-C) or 4 after AIA (Supplemental Figure 8B-D). Together, these findings suggest that FcyRI contributes to arthritis pain through a mechanism that parallels joint inflammation and cartilage damage in the AIA model.

Given the importance of neuronal FcyRI in regulating the excitability of primary sensory neurons (28, 29), we next asked whether neuronally-expressed FcyRI is involved in arthritis pain. In the AIA model, both male and female PirtCre; Fcgr1<sup>fl/fl</sup> mice subjected to AIA showed significantly reduced primary mechanical hyperalgesia in the ankle (Figure 8A; Supplemental Figure 9A) and secondary mechanical hyperalgesia in the hindpaw (Figure 8B,C; Supplemental Figure 9B,C) compared with PirtCre negative controls. However, there was no obvious difference between genotypes in either heat hyperalgesia in the hindpaw (Figure 8D; Supplemental Figure 9D) or joint inflammation following AIA (Figure 8E; Supplemental Figure 9E).

CFA-induced arthritis is another animal model of inflammatory arthritis (36-38), in which serum levels of specific rheumatoid and immunological biomarkers, such as rheumatoid factor and IgG, are elevated (39). In this model, deletion of Fcgr1 in sensory neurons significantly attenuated not only primary (Figure 8F) and secondary (Figure 8G-H) mechanical hyperalgesia, but also secondary thermal hyperalgesia in the hindpaw (Figure 8I). No significant difference in joint inflammation was observed between genotypes (Figure 8J). In addition, there was no difference between genotypes in the nocifensive behavior elicited by intraplantar injection of formalin, a short-term inflammatory agent (Figure 8K-L). These findings suggest that neuronal FcyRI is critical to the development and maintenance of arthritis pain but is apparently not required for joint inflammation in the AIA or CFA models. However, neuronal FcyRI is dispensable for some types of inflammatory pain.
Acute pharmacological blockade of peripheral FcγRI reverses arthritis pain in the AIA model

To circumvent potential confounding effects of genetic deletion of Fcgr1, we investigated whether acute pharmacological blockade of FcγRI at the periphery would attenuate arthritis pain. To test this possibility, we injected anti-CD64 monoclonal antibody (2.25 µg; 5 µl) or isotype control IgG2b into the inflamed ankle joint of WT mice once daily on days 1 and 2 after AIA (Figure 9A). In both male and female mice with established articular hypernociception, acute i.a injection of anti-CD64 antibody produced rapid reductions in mechanical hyperalgesia in the inflamed ankle and in both mechanical and thermal hyperalgesia in the ipsilateral hindpaw within 3 h, but did not attenuate joint swelling (Figure 9B-F; Supplemental Figure 10A-E). By contrast, i.a injection of isotype control IgG2b had no such effects. Next, we further asked whether this analgesic effect of anti-CD64 is secondary to a reduction in joint inflammation using joint histological H&E staining. Neutralizing peripheral FcγRI did not significantly affect immune cell infiltration in the AIA joint compared to isotype control IgG2b (Figure 9G-H). These findings suggest that local neutralization of FcγRI in the already inflamed joint reduces hyperalgesia through direct action on a neuronal target and/or via inhibition of ongoing local pronociceptive signaling, but not via the attenuation of inflammation.

Genetic deletion of Fcgr1 reduces AIA-induced hyperactivity of joint sensory afferents

Our recent study revealed that joint sensory afferents exhibited both abnormal hyperactivity and mechanical hypersensitivity in vivo following AIA (33). We therefore used an in vivo DRG recording preparation to determine whether deletion of Fcgr1 would reduce hyperactivity of joint sensory afferents during AIA (33). Extracellular electrophysiological recordings were obtained on day 1 after challenge from Dil-labeled mechanosensitive sensory neurons with a receptive field (RF) within the vehicle- or mBSA-treated ankle. In Fcgr1+/− mice, a total of 29 (8 C and 21 Aδ fibers) and 36 (16 C and 20 Aδ fibers) joint innervating DRG neurons were recorded from
vehicle- and mBSA-challenge animals, respectively. In global *Fcgr1*<sup>−/−</sup> mice, a total of 21 (5 C and 16 Aδ fibers) and 28 (15 C and 13 Aδ fibers) joint sensory neurons were recorded from the vehicle and AIA group, respectively. All neurons tested had conduction velocities (CV) within the ranges of C- (≤ 1.5 m/s) or Aδ -fibers (1.5-15 m/s). The mean CVs of C- or Aδ -fibers were similar between genotypes and treatments (Figure 10A,B). Consistent with previous findings (33), all the tested Dil-labeled joint sensory neurons from vehicle-treated *Fcgr1*<sup>+/+</sup> mice were silent in the absence of exogenous stimuli, with no detectable spontaneous activity (SA) (Figure 10C,D). In contrast, 10 (8 C and 2 Aδ fibers) of 36 (27.8%) joint sensory neurons in *Fcgr1*<sup>−/−</sup> mice with AIA exhibited SA. Similar to vehicle-treated *Fcgr1*<sup>+/+</sup> mice, no SA was observed in any of 21 recorded joint sensory neurons from vehicle-treated *Fcgr1*<sup>−/−</sup> mice (Figure 10C,D). Under AIA conditions, the incidence of SA was significantly lower in *Fcgr1*<sup>−/−</sup> mice (3 of 28 neurons; 10.7%; 2C and 2 Aδ fibers), compared to *Fcgr1*<sup>+/+</sup> mice (Figure 10C,D).

Since mechanical sensitization of joint sensory neurons likely represents a critical neuronal mechanism of mechanical hyperalgesia (33), we next asked whether FcγRI contributes to mechanical hypersensitivity of joint sensory neurons during AIA. To avoid the confounding effects of spontaneous firing, we exclusively focused on joint sensory afferents that did not exhibit SA. Under normal conditions, no mechanically evoked after-discharges were observed in either *Fcgr1*<sup>+/+</sup> or *Fcgr1*<sup>−/−</sup> mice. In the setting of AIA, 3 of 26 (11.5%; 2 C and 1 Aδ fibers) joint sensory neurons of *Fcgr1*<sup>+/+</sup> mice displayed after-discharges following punctate mechanical stimulation (10 or 20 mN; 2 s) of their RF whereas no mechanically evoked after-discharges occurred in 25 joint sensory neurons recorded from *Fcgr1*<sup>−/−</sup> mice (Figure 10E,F). We further compared mechanical sensitivity of joint sensory neurons that did not exhibit either SA or after-discharges between genotypes during AIA. Under arthritic conditions, the mean number of APs evoked by each mechanical force (5 mN to 40 mN) was significantly less in joint sensory neurons of *Fcgr1*<sup>−/−</sup> mice compared to those of *Fcgr1*<sup>+/+</sup> animals (Figure 10G,H). These results
indicate that FcγRI contributes to the sensitization of joint sensory neurons in the context of arthritis.

Discussion

Conventional wisdom has held that FcγRI, a common immune receptor for IgG-IC, is expressed exclusively in immune cells, where it plays a critical role in the regulation of various immune responses (13-16). However, we and others have previously reported evidence for the expression of FcγRI, but not FcγRII or FcγRIII, in a subpopulation of DRG neurons of rats and mice, respectively (27, 28). The results of our present study further challenge the traditional view of exclusive FcγRI function in immune cells by revealing that FcγRI is functionally and anatomically expressed in a subset of joint-innervating sensory neurons. Using ISH, we detected Fcgr1 mRNA expression in joint-innervating DRG neurons of all size categories, including both peptidergic and nonpeptidergic neurons. We also demonstrated that neuronal FcγRI activation is sufficient to drive joint sensory neuron firing, a conclusion supported by our in vivo and in vitro assays showing that IgG-IC can directly activate joint sensory neurons at their somata and peripheral terminals in a manner dependent on neuronal FcγRI.

A second advance achieved by this study is the revelation of a novel role for FcγRI, and for neuronally expressed FcγRI in particular, in modulating joint pain in both naïve and arthritis states. Acute pharmacological blockade, global deletion, and conditional deletion of Fcgr1 in primary sensory neurons each effectively alleviated the pronociceptive action of IgG-IC under naïve conditions and arthritis pain in the AIA and CFA models. Strikingly, these analgesic effects seem to be dissociable from effects on joint inflammation. Therefore, we suggest that peripheral FcγRI signaling, specifically in primary sensory neurons, contributes to arthritis pain through a mechanism that parallels inflammation and other pathological processes.

Sensitization of nociceptive joint sensory fibers likely represents a key mechanism in arthritis pain. In a recent study, we found that AIA enhances the in vitro excitability of joint innervating
DRG neurons (33). In addition, in vivo extracellular electrophysiological recordings on intact DRG showed that AIA causes spontaneous sensory neuronal activity and increases the responses of joint sensory neurons to mechanical stimulation (33). Enhanced mechanical sensitivity of joint sensory neurons might provide a peripheral neural basis for the behavioral signs of primary mechanical allodynia and/or hyperalgesia that accompany RA. Moreover, increased peripheral sensory afferent input might trigger ongoing spontaneous pain in arthritis and contribute to the development and maintenance of central sensitization. In the present study, we revealed that genetic deletion of Fcgr1 reduced both the incidence of abnormal activity of joint sensory fibers and mechanical hypersensitivity of joint sensory neurons after AIA, suggesting that FcγRI is necessary to sustain aberrant peripheral nociceptive activity in the context of this model of arthritis. Given that FcγRI is widely expressed in immune cells and is of critical importance to immune modulation, proinflammatory cytokines released within the damaged tissue under arthritic conditions might be important mediators of this process. However, our present study and reports from other groups suggest that FcγRI is not a major driver of joint inflammatory processes in the AIA model (18, 20). Therefore, it is plausible that FcγRI promotes AIA-induced hyperactivity of joint sensory afferents through a mechanism that operates in parallel with immune modulation. The expression and function of FcγRI in joint sensory afferents raise the possibility that neuronal FcγRI might be directly involved in this process. In our previous study, we demonstrated that neuronal FcγRI is functionally coupled to TRPC3 via the Syk-PLC-IP3 pathway to regulate the excitability of DRG neurons (29). Further work is therefore warranted to explore whether TRPC3 is involved in FcγRI-mediated arthritis pain. Neuronal FcγRI may also contribute to AIA-associated joint pain indirectly, by modulating the release of pain mediators such as CGRP and substance P from joint sensory neurons. Indeed, in cultured DRG neurons, FcγRI has been reported to mediate IgG-IC induced substance P release (27), which may in turn activate or sensitize joint sensory neurons through its own receptor in a paracrine or autocrine manner (40). This might account for AIA-induced
sensitization of joint sensory neurons that do not express FcγRI. Further investigation using sensory neuron subtype-specific Fcgr1 knockout mice will be necessary to assess these potential cell-autonomous and -nonautonomous roles of neuronal FcγRI.

RA is characterized by the accumulation of IgG-IC at the affected joint and the production of autoantibodies (41). Much attention has been focused on the role of IgG-IC/FcγRI signaling in RA pathogenesis (18, 35, 42, 43). Yet, little is known of its contribution to joint pain that accompanies this disorder. In this study, four orthogonal and complementary lines of evidence support the hypothesis that IgG-IC/FcγRI signaling, particularly in neurons, is necessary and sufficient to mediate joint pain via a mechanism that is at least to some extent dissociable from joint inflammation.

First, in naïve mice, exogenous application of IgG-IC elicited joint nociceptive behaviors whereas the pronociceptive effect of IgG-IC was diminished in mice lacking Fcgr1. It is conceivable that IgG-IC/FcγRI signaling could induce joint pain hypersensitivity by mobilizing immune cells and inflammatory responses. We did not, however, observe any obvious visual or histological signs of joint inflammation at early stages after IgG-IC injection. Moreover, qPCR analysis did not detect any acute alterations in the mRNA expression levels of various inflammatory mediators in the synovium following IgG-IC injection. In addition, depletion or deletion of immune cells had no impact on the pronociceptive action of IgG-IC. Based on the time scale of rapid modulation of neuronal activity by ligand (minutes to hours), the transient changes in pain behavior after i.a injection of IgG-IC are likely mediated by neuronal activation. The present findings, showing that IgG-IC had a direct action on neuronal activity in vivo and in vitro, thus support the view that IgG-IC may directly activate joint sensory neurons though FcγRI to elicit joint pain hypersensitivity in the naïve state. The involvement of neuronally expressed FcγRI was further validated by employing our newly generated conditional Fcgr1 knockout mice in which Fcgr1 is specifically deleted in primary sensory neurons. Conditional deletion of Fcgr1 remarkably suppressed IgG-IC-evoked nocifensive behaviors in naïve mice.
Second, FcγRI expression and function were upregulated within DRG in the AIA model. In RA patients, FcγRI is expressed de novo in the synovium (44) and its expression level in neutrophils, monocytes and synovial macrophages is upregulated (19, 45, 46). The mechanisms underlying the upregulation of FcγRI signaling require further study. It is probably due to the activation of a signaling cascade of inflammatory cytokines during arthritis, which upregulates FcγRI expression (47). In addition, cytokines may enhance the function of FcγRI by increasing the binding of IgG-IC to FcγRI (48). In this study, we revealed that a larger proportion of FcγRI-expressing joint sensory neurons displayed CGRP immunopositivity after AIA. Activation of neuronal FcγRI may result in more CGRP release from joint sensory afferents under arthritic conditions. The release of CGRP has been implicated in the generation of pain in certain arthritis models (49, 50). Therefore, it is possible that neuronal FcγRI contributes to arthritis pain through a mechanism involving neurogenic inflammation. However, deletion of neuronal Fcgr1 attenuated mechanical hyperalgesia but not joint swelling during AIA, suggesting that FcγRI expressed on peptidergic sensory neurons may signal hyperalgesia, but may not promote neuroinflammatory responses via neurogenic inflammation. Given that activation of sensory neurons is able to regulate innate and adaptive immunity (51), however, we cannot completely rule out a possible immune contribution of neuronal FcγRI to RA pain, especially in the late phase.

Third, global deletion of Fcgr1 attenuated arthritis pain in the AIA model. These inhibitory effects are unlikely secondary to the reductions in joint inflammation for several reasons. Previous studies have demonstrated that FcyRI is not essential for inflammatory processes in this model (18, 20). Moreover, in line with previous studies (52), our qPCR assay revealed an upregulation of inflammatory mediators in the inflamed synovium of mice with AIA. Yet, the extent of the upregulation was similar between WT and global Fcgr1−/− mice. IHC staining and joint histological analysis further showed that immune cell infiltration in the AIA joint was not different between genotypes. Although FcγRI has been implicated in cartilage destruction in the
AIA model (18, 35), we did not observe obvious histological changes in cartilage in the inflamed joints at the early phase of AIA (i.e. day 2 and 4). One possible explanation for this discrepancy is that obvious cartilage destruction may only occur at the later time points (i.e., day 7) in the AIA model (18). Thus, our findings argue against the possibility that global deletion of Fcgr1 attenuated arthritis pain primarily via reductions of cartilage destruction.

Fourth, acute blockade of FcγRI at the periphery, achieved with local injection of anti-CD64 antibody, markedly attenuated established arthritis pain, but not joint swelling, in the AIA model. Joint histological analysis further confirmed that acute pharmacological inhibition of FcγRI had no detectable impact on immune cell infiltration into the inflamed synovium. It is therefore likely that the analgesic effect of neutralizing FcγRI is due to direct action on a neuronal target and/or inhibition of ongoing local pronociceptive signaling, and not to the attenuation of joint inflammation.

Although RA pain is often thought to be of inflammatory origin, joint inflammation alone does not entirely account for arthritis pain (5, 6). The present study sheds light on a novel mechanism that might contribute to RA-associated pain. We suggest that IgG-IC accumulated in the inflamed joint is sufficient to directly activate and sensitize joint sensory neurons through neuronal FcγRI to evoke joint pain. Since no single experimental model of arthritis recapitulates all aspects of human RA (53), our data from the AIA and CFA models might not be completely generalizable to other models. We also cannot exclude the contributions of neuronal FcγRI to inflammation-related events beyond the sensitivity of our histological and biochemical assays. In addition, our data do not rule out the involvement of other central and peripheral non-inflammatory mechanisms in arthritis pain. Nevertheless, this proposed new location and role for IgG-IC/FcγRI signaling might be especially relevant to joint pain that occurs prior to the detectable onset of inflammation or that which persists even after apparently successful reversal of joint inflammation. Critical assessment of neuronal FcγRI contributions to additional arthritis models and future translational studies are therefore warranted to define the biological
relevance of our findings, and to determine whether neuronal FcγRI merits consideration as a therapeutic target for the treatment of RA pain.

Methods

Animals

Animals were housed under a 12-hour light/dark cycle with ad libitum access to food and water. Male mice used in the study were 2 to 3 months of age and weighed 20-30 g. In some experiments, female mice were also used as indicated. Breeders of T or B cell-deficient Rag1\(^{-/-}\) mice, mast cell-deficient c-Kit\(^{W-sh/W-sh}\) mice were obtained from Jackson Laboratories (Bar Harbor, ME). Breeders of global Fcgr1\(^{-/-}\) mice and PirtCre-GCamp6 mice were generously provided by Dr. Sjef Verbeek (Leiden University Medical Center, The Netherlands) and Dr. Xinzhong Dong (Johns Hopkins University), respectively. Littermates were generated by interbreeding heterozygotes on the C57BL/6 background.

Generation of Fcgr1\(^{fl/fl}\) mice

Fcgr1\(^{fl/fl}\) mice were generated using CRISPR/Cas9 genome editing. Two loci in the Fcgr1 gene were targeted for loxP site insertion, one 283 nucleotides upstream of exon 1 and another 223 nucleotides downstream of exon 3, >200 bp away from splice donor or acceptor sequences. The loxP sites were designed close enough to one another (<2.5kb) to allow efficient Cre-mediated recombination. Guide RNA (gRNA) sequences were designed using the CRISPR Design Tool (CRISPR.mit.edu), to target these loci within the Fcgr1 gene while minimizing off-target mutations. crRNAs corresponding to each gRNA (5'- CAG UAA ACC CUG AAA GAG UGG UUU UAG AGC UAU GCU GUU UUG -3' for the gRNA upstream of Exon 1 and 5'- AUC CCC CGC UGG ACC AGU UUG UUU UAG AGC UAG GCU GUU UUG-3' for the gRNA downstream of exon 3) were synthesized commercially (Dharmacon, Lafayette, CO). Two corresponding homology repair templates (HRT) containing a loxP site with adjacent restriction
sites and flanked on each side by 36 nucleotides of target sequence homology were generated by polymerase chain reaction (PCR) (54), using a loxP site-containing plasmid (pBS-loxp plasmid, gift of Dr. Randall Reed, Johns Hopkins University) as a template and the following primers: HRT upstream of exon 1: forward: 5'-ATT CAG GCT ATC AGA GCT ACA GTA AAC CCT GAA AGA GCG GTG GCG GCC GCT CTA GA-3'; reverse: 5'-GCT ACC ATG ACT AGC TAC ATA TCC CTC CCA CCT CAC TCC TGC AGC CCG GGG GAT CC-3'; loxp insertion at the downstream of exon 3: forward: 5'-AGG CCC TAT TTG CCT GCA GCA TCC CCC GCT GGA CCA GCG GTG GCC GCT CTA GA-3'; reverse: 5'-GAG CCC GGG ATT TTT GGG TGA CAC TGT CAC CAA AAC TCC TGC AGC CCG GGG GAT CC-3'.

crRNAs, trRNA, Cas9 protein (provided by JHU Transgenic Mouse Core Facility), and the double-stranded HRT DNAs were microinjected into C57BL6 mouse zygotes and implanted into pseudopregnant females by the Johns Hopkins School of Medicine Transgenic Mouse Core Facility. Tail genomic DNA from founders and offspring was genotyped using two sets of primers [for loxp insertion at 5' UTR of exon1, forward (UF): 5'- GAT CTC TGT GAG GTC AAG GCT-3'; reverse (UR): 5'-CCT CCC AAG TGC TAG GAT TAT-3'; for loxp insertion at the downstream of exon 3, forward (3-4F): 5'-GTC AAA TCA GGT CAG ACA GCT-3'; reverse (3-4R): 5'-AGA ACT GCT GTG GGT GAA GCT-3'], as shown in Figure. 5A. Only one of 26 founders possessed two loxP sites inserted to the same allele. This founder was then backcrossed 10 generations against C57BL6 mice to eliminate off-target mutations. All the offspring developed normally except that about 10 of them obtained during the first 3-5 generations of backcross exhibited spontaneous turning behavior and were excluded from behavior studies. To generate sensory neuron-specific Fcgr1 knockout mice (PirtCre; Fcgr1\textsuperscript{fl/fl}), Fcgr1\textsuperscript{fl/fl} mice were crossed onto PirtCre mice to achieve Fcgr1 deletion in peripheral sensory neurons. The efficiency of Fcgr1 mRNA deletion in DRG was confirmed by quantitative real-time PCR using one set of primers [Figure 5A; forward (1F): 5'-CAG CCT CCA TGG GTC AGT-3'; reverse (3R): 5'-TGA AAA CTG GCC TCT GGG AT-3'].
Preparation of IgG-immune complex (IgG-IC)

IgG-IC was formed by incubating BSA (10 μg/ml; Sigma St. Louis, MO) and anti-BSA IgG (5 mg/ml; MP Biomedical, Solon, OH) at the mass ratio of 1:2 for 1 h at 37°C as described previously (55). The IgG-IC was then diluted to the concentrations of 1, 10 and 100 μg/ml. Given that the estimated molecular mass of immune complexes is typically around 650-850 kDa (56), the molar concentration of 1 μg/ml is equivalent to 1.2-1.5 nM. Doses were chosen based on our pilot studies and other reports (57).

Intra-articular injection of IgG-IC and anti-FcγRI (CD64) antibody

IgG-IC (1, 10, 100 μg/ml; 10 μl) was injected into right knee (for joint histology and real-time PCR on synovium tissue) or ankle (for behavioral testing) joint cavity of naïve mice. The same amounts of monomeric IgG and PBS served as controls. Pain-related behaviors and ankle diameter were measured 1 – 24 h after injection. Since the concentration of 100 μg/ml is within the range of IgG-IC accumulated in the serum in RA patients (57), this concentration was chosen throughout this study. In some experiments, either rat anti-mouse FcγRI monoclonal antibody (2.25 μg in 10 μl saline; R&D system, Minneapolis, MN), or isotype control (monoclonal rat IgG2b; R&D system, Minneapolis, MN) was injected into the right knee (for joint histology) or ankle (for behavioral testing) of mice on days 1 and 2 after AIA. Pain-related behaviors and joint swelling were assessed over the ensuing 3 h.

Retrograde labeling of ankle joint sensory afferents

For in vivo and in vitro studies, DRG cell bodies with their afferent fibers innervating ankle joints were identified by the presence of a retrogradely transported red fluorescent dye, Dil (Sigma, St. Louis, MO). Dil was injected into the right ankle (2.5 mg/ml, 8 μl in 25% ethanol) at least 1 week before harvesting. Since our pilot study showed that Dil was not compatible with in situ
hybridization assay, fast blue (FB; 8 µl; 1% in saline; Polysciences Inc, Warrington, PA) was alternatively injected to the ankle cavity at least 2 weeks before harvesting.

**In situ hybridization (ISH)**

SP6 transcribed antisense and T7 transcribed sense control probes were synthetized from mouse *Fcgr1* (NM_010186) cDNA clone (MR225268, OriGene) using one set of primers (forward: 5’-ATT TAG GTG ACA CTA TAG AAT CCT CAA TGC CAA GTG ACC C-3’; reverse: 5’-GCG TAA TAC GAC TCA CTA TAG GGC GCC ATC GCT TCT AAC TTG C-3’). The antisense probe was designed to target exons 1-6 of the mouse *Fcgr1* gene. The probes were then labeled using a digoxygenin (DIG) RNA labeling kit (Roche 11277073910) according to the manufacturer’s instructions. Prehybridization, hybridization and washing were performed on DRG and spleen sections using standard methods (58). The temperatures for prehybridization/hybridization and for washing were 60°C and 62°C, respectively. To combine ISH with IHC, tissue sections were incubated with sheep anti-DIG antibody (Table S2) and subjected to the standard IHC protocol as above.

**Knee histology**

Hind knee joints from mice were post-fixed in 4% PFA for 48 h, decalcified in 10% EDTA (Sigma) for 3-4 weeks, then dehydrated in ethanol and embedded in paraffin. Sections (5 µm) were cut and stained with hematoxylin & eosin (H&E), or toluidine blue (TB)/fast green, and scored by the evaluators in a blinded manner. Three sections per knee joint at different depths were analyzed for synovitis on a scale from 0 (normal) to 3 (severe) (59) and for cartilage damage on a scale from 0 (normal) to 6 (bone loss, remodeling, deformation) (60), respectively. The score per knee was averaged.

**Statistics**
Data were presented as means ± SEM. Two-tailed Student’s t-test was used to test the significance of differences between two groups. Comparisons for multiple groups or multiple time points were carried out using a one-way or two-way analysis of variance (ANOVA) for random measures or repeated measures followed by Bonferroni’s or Tukey’s post hoc test comparisons. Comparisons of proportions were made by $\chi^2$ test. P less than or equal to 0.05 was considered significant. Group sizes and the type of statistical tests used for each comparison were noted in each figure legend.

**Study approval**

All animal experimental procedures were approved by the Institutional Animal Care and Use Committee of Johns Hopkins University School of Medicine and were in accordance with the guidelines provided by the National Institute of Health and the International Association for the Study of Pain.
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Author contributions

L.Q. conceived the project, designed the experiments, conducted behavioral tests, in vivo and in vitro calcium imaging, and in vivo electrophysiological experiments, analyzed the data, and wrote the manuscript; L.W. performed IHC and joint histology staining, quantitative RT-PCR and in situ hybridization experiments; X. J. and S.M. J. carried out quantitative RT-PCR; Q.Z. carried out in vivo calcium imaging; T.C. performed joint histology staining; Y.L. carried out behavioral tests; R.R. and H.K. assisted with gRNA and HDR vector design and founder genotyping; M.J.C and X.D. facilitated experimental design and analysis and revised the manuscript.
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Fig. 1

A FB Fcgr1 mRNA NeuN Merge

B Fcgr1 mRNA Fcgr1 mRNA/NeuN

C Sense control Sense control/NeuN

D FB Fcgr1 mRNA Peripherin Merge

FB Fcgr1 mRNA NF200 Merge

FB Fcgr1 mRNA CGRP Merge

FB Fcgr1 mRNA GS Merge

% of Fcgr1-labeled joint sensory neurons
Figure 1. Analysis of FcγRI expression in mouse joint sensory neurons. Joint innervating DRG neurons were labeled retrogradely by injection of fast blue (FB; 8 μl, 1% in saline) into the ankle cavity at least 2 weeks before harvesting. (A) In situ hybridization (ISH) images showing Fcgr1 mRNA expression (green) in a subset of FB-labeled joint sensory neurons (blue) of WT mice (n = 5 mice). Fcgr1 signal was colocalized with the pan-neuronal marker, NeuN (red). (B) ISH image showing absence of Fcgr1 mRNA expression in DRG neurons of global Fcgr1−/− mice (n = 3 mice). (C) ISH image of sense control probe (n = 3 mice). (D) Fluorescent ISH for Fcgr1 and immunostaining using antibodies against peripherin (n = 5 mice), NF200 (n = 5 mice), CGRP (n = 5 mice), and glutamine synthetase (GS; n = 6 mice), along with quantitative analysis of percentage overlap. Scale bar, 50 μm.
Fig. 2

A

Fcgr1+/+

Fcgr1−/−

Ag IgG-IC CAP

Time (s)

Ratio (340/380)

0 0.5 1 1.5 2

0 0.5 1 1.5 2

Time (s)

B

% of joint sensory neurons

Ag Ab IgG-IC IgG-IC

(75/427)

(29/427)

(5/130)

C

Dil

PBS

Pinch

Fcg1+/+

Fcg1−/−

Dil

Monomeric IgG

Pinch

Dil

IgG-IC

Pinch

Dil

IgG-IC

Pinch

Dil

IgG-IC

Pinch

D

% of responsive joint sensory neurons

PBS IgG IgG-IC IgG-IC

Fcg1+/+

Fcg1−/−
Figure 2. IgG-IC directly activates joint sensory neurons through FcγRI in vitro and in vivo. (A) Representative traces of Ca^{2+} responses evoked by antigen (Ag) alone (BSA), IgG-IC (1 µg/ml, 2 min) and capsaicin (CAP; 1 µM,10 s) in joint innervating (Dil labeled) DRG neurons from Fcgr1^{+/-} (left) and global Fcgr1^{-/-} (right) mice. (B) Quantitative analysis showed that IgG-IC, but not Ag or antibody (Ab; anti-BSA) alone, evoked Ca^{2+} response in a larger proportion of Dil-labeled joint sensory neurons from Fcgr1^{+/-} mice than those from Fcgr1^{-/-} mice. *p < 0.05 versus Fcgr1^{+/-}, # p < 0.05 versus Ag, χ^2 test. Number of responsive neurons and total number tested are in parentheses. (C) Left, representative Dil fluorescence (red) in L4 DRG neuronal cell bodies in PirtCre-GCaMP6 mice that were either Fcgr1^{+/-} or Fcgr1^{-/-}, retrogradely labeled with ankle joint injection of Dil (2 mg/ml, 8 µl in saline). Right, GCAMP6 fluorescence (green) in the same fields before and after stimulation of the RF with the indicated stimuli. White arrows show DRG neurons from Fcgr1^{+/-} mice exhibiting an increase in GCaMP6 fluorescence when the ankle was squeezed with blunt forceps and 4 min after IgG-IC (100 µg/ml; 10 µl) was injected into ankle joint cavity, but not after vehicle (PBS; 10 µl) or monomeric IgG (100 µg/ml; 10 µl) was injected. By contrast, little or no increase in GCaMP6 fluorescence increase was induced by IgG-IC in joint sensory neurons from Fcgr1^{-/-} mice. Scale bar: 50 µm. (D) Quantitative analysis of Ca^{2+} responses to PBS (n = 6 mice), monomeric IgG (n = 6 mice), IgG-IC (n = 10 mice) in joint sensory neurons of Fcgr1^{+/-} mice and IgG-IC in Fcgr1^{-/-} mice (n = 8 mice). *p < 0.05 versus Fcgr1^{+/-}, # p < 0.01 versus PBS; one-way ANOVA followed by Tukey's test.
**Fig. 3**

A. Ankle mechanical threshold (g) with hours after injection.

B. PWF (%) with hours after injection.

C. PWF (%) with hours after injection.

D. PWL (s) with hours after injection.

E. Joint diameter (mm) with hours after injection.

F. Immunofluorescence images showing Ly6C/G/DAPI, CD68/DAPI, and c-Kit/DAPI.

G. Fluorescence intensity (A.U.) for Ly6C/G and CD68.

H. Synovitis score for PBS, IgG, and IgG-IC.
Figure 3. IgG-IC elicits acute articular hypernociception in naïve mice. (A-E) Mice were injected intra-articularly (i.a.) with IgG-IC (1, 10, 100 µg/ml; 10 µl), monomeric IgG (100 µg/ml; 10 µl), or vehicle (PBS; 10 µl), and pain-like behaviors and joint diameter were evaluated over 1-24 h. Injection of IgG-IC, but not monomeric IgG, reduced mechanical threshold in the ankle (A) and increased the frequency of paw withdrawal in response to application of 0.07 and 0.4 g force via a von Frey filament (B-C), but did not induce heat hyperalgesia (D) or visible joint swelling (E) in the ipsilateral paw, compared to vehicle. n = 8-10 mice /group; * p < 0.05 versus PBS; #p< 0.05 versus before injection; two-way ANOVA for repeated measures followed by Bonferroni post hoc test. PWF: Paw Withdrawal Frequency; PWL: Paw Withdrawal Latency. (F) Representative sections of knee joints taken 1 h after i.a injection with either PBS, monomeric IgG, or IgG-IC and stained for Ly6C/G, CD68, CD3 or c-Kit. s: synovium. Scale bar: 200 µm. (G) Quantification showed no significant differences between treatment groups. n = 4-5 mice/group; p > 0.05; one-way ANOVA followed by Tukey’s test. (H) Representative sections of knee joint taken 1 h after i.a injection with PBS, monomeric IgG or IgG-IC, stained with H&E and scored for synovitis. s: synovium. Scale bar: 100 µm. No significant difference in synovitis score was observed between treatments. n = 3-4 mice/group; p > 0.05; one-way ANOVA followed by Tukey’s test.
Fig. 4

(A) Baseline

Ankle mechanical threshold (g)

Fcgr1+/+  Fcgr1−/−

0 200 400

(B) Baseline

Paw withdrawal frequency (%)

Fcgr1+/+  Fcgr1−/−

0 0.5 1.0 1.5

Von Frey force (g)

(C) Baseline

Paw withdrawal latency (s)

Fcgr1+/+  Fcgr1−/−

0 5 10 15

(D) Baseline

Ankle mechanical threshold (g)

Fcgr1+/+  Fcgr1−/−

-10 1 2 3 4 5 22 24

Hours after injection

(E) Baseline

PWF to 0.07 g force (%)

Fcgr1+/+  Fcgr1−/−

-10 1 2 3 4 5 22 24

Hours after injection

(F) Baseline

PWF to 0.4 g force (%)

Fcgr1+/+  Fcgr1−/−

-10 1 2 3 4 5 22 24

Hours after injection

(G) Baseline

Ankle mechanical threshold (g)

Veh  Clodronate

-10 1 2 3 4 5 22 24

Hours after injection

(H) Baseline

PWF to 0.07 g force (%)

Veh  Clodronate

-10 1 2 3 4 5 22 24

Hours after injection

(I) Baseline

Ankle mechanical threshold (g)

Rag1+/+  Rag1−/−

-10 1 2 3 4 5 22 24

Hours after injection

(J) Baseline

PWF to 0.07 g force (%)

Rag1+/+  Rag1−/−

-10 1 2 3 4 5 22 24

Hours after injection

(K) Baseline

Ankle mechanical threshold (g)

c-Kit+/+  c-Kit+/+ c-Kit+/+ c-Kit−/−

-10 1 2 3 4 5 22 24

Hours after injection

(L) Baseline

PWF to 0.07 g force (%)

c-Kit+/+  c-Kit+/+ c-Kit+/+ c-Kit−/−

-10 1 2 3 4 5 22 24

Hours after injection
Figure 4. FcγRI mediates IgG-IC-induced acute joint nocifensive behaviors in mice. (A-C) Comparison of basal mechanical sensitivity to ankle press (A) and to plantar stimulation with von Frey filaments (B), and basal thermal sensitivity to plantar application of radiant heat (C) between Fcgr1+/+ (n = 9 mice) and global Fcgr1−/− mice (n = 12 mice). p > 0.05; unpaired student’s t-test or two-way ANOVA for repeated measures followed by Bonferroni post hoc test. (D-F) Global Fcgr1−/− mice (n = 12 mice) exhibited a higher mechanical threshold in the ankle (D) and lower paw withdrawal frequency (PWF) to 0.07 (E) and 0.4 g force (F) applied to the hindpaw following i.a injection of IgG-IC (100 µg/ml; 10 µl) compared to Fcgr1+/+ control littermates (n = 9 mice). *p < 0.05 versus Fcgr1+/+ controls, #p < 0.05 versus before injection, two-way ANOVA for repeated measures followed by Bonferroni post hoc test. (G-H) Depletion of synovial macrophages with liposomal clodronate (5 mg/ml; 6 µl) had no significant effects on mechanical hyperalgesia in the ankle (G) or hindpaw (H) in mice upon injection of IgG-IC, compared to liposomal control (Veh). n = 10 mice/group; p > 0.05, two-way ANOVA for repeated measures followed by Bonferroni post hoc test. (I-L) No significant differences were seen in IgG-IC induced mechanical hyperalgesia in the ankle and hindpaw in mice lacking T cells (Rag-1−/−) or mast cells (c-Kit W−/−), compared with WT controls. n = 10-11 mice/group; p > 0.05, two-way ANOVA for repeated measures followed by Bonferroni post hoc test.
**Figure 5.** Neuronal FcγRI contributes to IgG-IC-induced acute nocifensive behaviors. (A) Strategy for generation of primary sensory neuron selective FcγRI knockout mice. Two loxp sites were inserted 5’ to exon 1 and 3’ to exon 3 of the FcγRI gene, respectively. Primers UF and UR, 3-4F and 3-4R, respectively, were used to confirm correct loxp insertions at each site. Deletion of FcγRI gene in primary sensory neurons was achieved by crossing FcγRIfl/fl mice with PirtCre mice. (B) qRT-PCR analysis using primers 1F and 3R from panel (A) revealed a significant reduction in FcγRI mRNA expression in DRG tissue (n= 10-11 mice/group) but not in spleen of PirtCre; FcγRIfl/fl mice (n = 4-5 mice/group) compared to FcγRIfl/fl controls. (C) Representative ISH image on DRG and spleen. Scale bar, 50 µm. Inset shows area of high power magnification. Scale bar, 20 µm. Quantification shows reductions in FcγRI mRNA expression in DRG neurons (NeuN) but not in spleen macrophages (F4/80) of PirtCre; FcγRIfl/fl mice compared to FcγRIfl/fl controls. n = 4 mice/group; *p < 0.05 versus FcγRIfl/fl controls. For panel B and C, unpaired Student’s t test were used. (D-F) No significant differences were observed between genotypes in basal mechanical sensitivity in the ankle (D) or hindpaw (E), or in basal thermal sensitivity in the hindpaw (F). n = 10-19 mice/group, p >0.05, unpaired Student’s t test or two-way ANOVA for repeated measures followed by Bonferroni post hoc test. (G-I) Time course of mechanical threshold in the ankle and paw withdrawal frequency (PWF) to 0.07 and 0.4 g force before and after i.a. injection of IgG-IC (100 µg/ml; 10 µl). n = 9 mice/group; *p< 0.05 versus FcγRIfl/fl controls; #p < 0.05, ## p < 0.01 versus before injection; two-way ANOVA for repeated measures followed by Bonferroni post hoc test.
Fig. 6

A) Day 1

B) Day 3

C) Ctrl | AIA

FB

Fcgr1 mRNA

CGRP

NeuN

Merge

D)

% of Fcgr1* neurons in all DRG neurons

E)

% of Fcgr1* neurons in FB-labeled joint sensory neurons

F)

% of CGRP+ neurons in Fcgr1* joint sensory neurons

G)

% of IgG+ responsive joint sensory neurons

75/427

47/177

75/427

47/177

**

*
Figure 6. AIA upregulates the expression and function of FcγRI in mouse DRG neurons. (A-B) qRT-PCR analysis of Fcgr1 mRNA expression in the DRG of control (Ctrl) and AIA mice on days 1 (A; n = 8 mice/group) and 3 (B; n = 5 mice/group) after challenge. *p < 0.05 versus control; unpaired Student’s t test. (C) Representative lumbar DRG ISH for Fcgr1, and immunostaining for peripherin, NF200 and CGRP, and the merged image from control (Ctrl; n = 6) and AIA (n = 6) mice. DRG neurons innervating ankle joint were retrogradely labeled with FB. Scale bar, 50 µm. (D) Percentage of Fcgr1+ neurons among all DRG neurons in control and AIA mice. (E) Percentage of Fcgr1+ joint sensory neurons in control and AIA mice. (F) Percentage of CGRP+ neurons among Fcgr1+ joint sensory neurons in control and AIA mice. n = 6 mice/group; *p < 0.05 versus control, unpaired Student’s t test. (G) Quantitative analysis of Fura-2 Ca2+ imaging shows that a larger proportion of Dil-labeled joint sensory neurons from AIA mice responded to IgG-IC (1 µg/ml, 2 min) compared to those from control mice. *p < 0.05 versus control, χ2 test. Numbers of responsive neurons divided by total number tested are noted above graphs.
Fig. 7

A  Ankle mechanical threshold (g)

B  0.07 g force

C  0.4 g force

D  PWL (s)

E  Joint diameter (mm)

- Fcgr1^{+/+} Ctrl
- Fcgr1^{+/+} AIA
- Fcgr1^{-/-} Ctrl
- Fcgr1^{-/-} AIA

Days after challenge
**Figure 7.** FcγRI modulates arthritis pain in the AIA model. (A) Time course of mechanical threshold in the ankle, paw withdrawal frequency (PWF) in response to 0.07 g and 0.4 g force in the hindpaw, paw withdrawal latency (PWL) to radiant heat in the hindpaw, and ankle joint diameter in \( Fcgr1^{+/+} \) and global \( Fcgr1^{-/-} \) mice after vehicle control (Ctrl) and mBSA (AIA) challenge. \( n = 7-11 \) mice/group; \( *p < 0.05 \) versus \( Fcgr1^{+/+} \); \( #p < 0.05 \) versus day 0; two-way repeated measures ANOVA followed by Bonferroni post hoc test.
Fig. 8

A. Ankle mechanical threshold (g) over days after AIA.

B. Percent PWF to 0.07 g force (%) over days after AIA.

C. Percent PWF to 0.4 g force (%) over days after AIA.

D. Paw withdrawal latency (s) over days after AIA.

E. Joint diameter (mm) over days after AIA.

F. Ankle mechanical threshold (g) over weeks after CFA injection.

G. Percent PWF to 0.07 g force (%) over weeks after CFA injection.

H. Percent PWF to 0.4 g force (%) over weeks after CFA injection.

I. Paw withdrawal latency (s) over weeks after CFA injection.

J. Joint diameter (mm) over weeks after CFA injection.

K. Nociceptive behavior duration (s) after formalin injection.

L. Nociceptive behavior duration (s) after formalin injection.
Figure 8. Neuronal FcyRI mediates arthritis pain in the AIA and CFA models. (A-E) Time course of ankle mechanical threshold (A), paw withdrawal frequency (PWF) to 0.07 g (B) and 0.4 g force (C), paw withdrawal latency to radiant heat (D), and ankle joint diameter (E) of male mice following AIA in PirtCre; Fcgr1<sup>fl/fl</sup> mice and Fcgr1<sup>fl/fl</sup> control littermates. n = 10 mice/group; *p < 0.05 versus Fcgr1<sup>fl/fl</sup> controls, #p < 0.05 versus day 0, two-way repeated measures ANOVA followed by Bonferroni post hoc test. (F-J) Ankle mechanical threshold (F), paw withdrawal frequency (PWF) to 0.07 g (G) and 0.4 g force (H), paw withdrawal latency to radiant heat (I), and ankle joint diameter (J) in male Fcgr1<sup>fl/fl</sup> and PirtCre; Fcgr1<sup>fl/fl</sup> mice after injection CFA (5 µl) into the ankle joint. n = 10 mice/group; *p < 0.05 versus Fcgr1<sup>fl/fl</sup> controls, #p < 0.05 versus week 0, two-way repeated measures ANOVA followed by Bonferroni post hoc test. (K-L) Total duration of pain-related behavior during the 1<sup>st</sup> (1-10 min) and the 2<sup>nd</sup> phases (11-60 min) after intraplantar injection of formalin (5%; 20 ul) in Fcgr1<sup>fl/fl</sup> and PirtCre; Fcgr1<sup>fl/fl</sup> mice (both genders). n = 8 mice/group. p > 0.05 versus Fcgr1<sup>fl/fl</sup> controls, one-way repeated measures ANOVA followed by Bonferroni post hoc test.
Fig. 9

A. Day 1 and 2 after AIA

i.a. Anti-CD64 IgG2b isotype

Behavioral testing or tissue harvesting

B. Ankle mechanical threshold (g)

C. PWF to 0.07 g force (%)

Days after AIA

D. PWF to 0.4 g force (%)

Days after AIA

E. Paw withdrawal latency (s)

Days after AIA

F. Joint diameter (mm)

Days after AIA

G. Control IgG2b

Anti-CD64

H. Synovitis score

Synovitis score

s
**Figure 9.** Acute pharmacological blockade of FcγRI attenuates AIA-associated pain in male mice. (A) Experimental schematic indicating once daily injection of anti-CD64 (2.25 µg; 5 µl) or IgG2b isotype control (2.25 µg; 5 µl) into knee (for histology) or ankle cavity (for behavioral testing) of mice on days 1 and 2 after AIA. Pain-related behaviors were measured within 3 h after each injection. (B-F) Effects of repeated daily i.a injection of anti-CD64 or IgG2b isotype control on mechanical threshold in the ankle (B), paw withdrawal frequency (PWF) in response to 0.07 g (C) and 0.4 g force (D), paw withdraw latency to radiate heat (E), and ankle joint diameter (F) in the mice with AIA. n = 9 mice/group; *p< 0.05 versus control IgG2b isotype; #p < 0.05, versus day 0; two-way ANOVA for repeated measures followed by Bonferroni post hoc test. (G) H&E staining assessment of synovitis in the inflamed knee joint of AIA mice treated with IgG2b isotype control (n = 5 mice) or anti-CD64 (n = 6 mice). s: synovium. Scale bar, 100 µm. (H) Quantification showed no difference in synovitis score between treatment groups. p > 0.05, unpaired Student’s t test.
**Fig. 10**

**A**
- CV of C fibers (m/s) for various conditions:
  - Ctrl
  - AIA

**B**
- CV of Aδ fibers (m/s) for various conditions:
  - Ctrl
  - AIA

**C**
- Afterdischarge (SA) for various conditions:
  - Ctrl
  - AIA

**D**
- SA incidence (%) for various conditions:
  - Ctrl
  - AIA

**E**
- Afterdischarge incidence (%)
  - Ctrl
  - AIA

**F**
- Afterdischarge incidence (%)
  - Ctrl
  - AIA

**G**
- Mechanical sensitivity for various conditions:
  - Ctrl
  - AIA

**H**
- Number of APs for various mechanical forces and conditions:
  - FcγRI<sup>+/+</sup>; Ctrl
  - FcγRI<sup>-/-</sup>; Ctrl
  - FcγRI<sup>+/+</sup>; AIA
  - FcγRI<sup>-/-</sup>; AIA
**Figure 10.** FcγRI contributes to hyperactivity of joint sensory neurons following AIA. (A-B) Distribution of the recorded C (A) and Aδ (B) fibers innervating the ankle of Fcgr1+/+ and Fcgr1−/− mice 1 day after vehicle control (Ctrl) or mBSA (AIA) challenge. No significant difference in CV was seen between treatments or genotypes, p > 0.05, two-way ANOVA followed by Bonferroni post hoc test. (C) Representative traces of abnormal spontaneous activity (SA) were recorded in Dil-labeled joint sensory neurons of mice. (D) Global Fcgr1−/− mice exhibited lower incidence of SA at day 1 after AIA, *p < 0.05 versus controls, #p < 0.05 versus Fcgr1+/+ mice, χ² test. Number of neurons tested is noted above graphs. (E) Responses of joint sensory neurons in Fcgr1+/+ and Fcgr1−/− mice to a 2 s, 10 mN mechanical stimulus delivered via a 100 μm probe in control (Ctrl) and AIA mice. (F) Prevalence of mechanically evoked after-discharges in joint sensory neurons of Fcgr1+/+ and Fcgr1−/− mice on day 1 after challenge, *p < 0.05 versus controls; #p < 0.05 versus Fcgr1+/+ mice; χ² test. Number of neurons tested is noted above graphs. (G) Representative responses of joint sensory neurons in Fcgr1+/+ and Fcgr1−/− mice to mechanical stimulation (2 s in duration) of their RF with von Frey filaments (100 μm tip diameter) at the indicated bending forces on day 1 after challenge. (H) The mean number of APs evoked by mechanical stimuli in joint sensory neurons from Fcgr1+/+ (Ctrl: 29 neurons; AIA: 23 neurons) and Fcgr1−/− mice (Ctrl: 21 neurons; AIA: 25 neurons) on day 1 after AIA, *p < 0.05, **p < 0.01 versus controls, #p < 0.05; #p < 0.01 versus Fcgr1+/+, two-way repeated measures ANOVA followed by Bonferroni post hoc test.