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Lysosomal β-glucuronidase regulates Lyme and rheumatoid arthritis severity

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Lyme disease, caused by the spirochete *Borrelia burgdorferi*, is the most prevalent arthropod-borne illness in the United States and remains a clinical and social challenge. The spectrum of disease severity among infected patients suggests that host genetics contribute to pathogenic outcomes, particularly in patients who develop arthritis. Using a forward genetics approach, we identified the lysosomal enzyme β-glucuronidase (GUSB), a member of a large family of coregulated lysosomal enzymes, as a key regulator of Lyme-associated arthritis severity. Severely arthritic C3H mice possessed a naturally occurring hypomorphic allele, *Gusb*<sup>b</sup>. C57BL/6 mice congenic for the C3H *Gusb* allele were prone to increased Lyme-associated arthritis severity. Radiation chimera experiments revealed that resident joint cells drive arthritis susceptibility. C3H mice expressing WT *Gusb* as a transgene were protected from severe Lyme arthritis. Importantly, the *Gusb*<sup>b</sup> allele also exacerbated disease in a serum transfer model of rheumatoid arthritis. A known GUSB function is the prevention of lysosomal accumulation of glycosaminoglycans (GAGs). Development of Lyme and rheumatoid arthritis in *Gusb*<sup>b</sup>-expressing mice was associated with heightened accumulation of GAGs in joint tissue. We propose that GUSB modulates arthritis pathogenesis by preventing accumulation of proinflammatory GAGs within inflamed joint tissue, a trait that may be shared by other lysosomal exoglucosidases.

Introduction

Lyme disease, caused by the spirochete *Borrelia burgdorferi* (1), is the most prevalent arthropod-borne illness in the United States. More than 30,000 cases are reported each year, while estimates suggest that around 300,000 are diagnosed annually (2, 3). Disease severity varies greatly among the affected population, with up to 60% of untreated patients developing a self-limiting, inflammatory arthritis (4, 5). Even following appropriate antibiotic therapy, up to 10% of patients will develop chronic arthritis, which can persist for months to years. While the genetic composition of the spirochete is a critical determinant of the invasive potential of the bacteria (6), other host-associated attributes are also clearly instrumental in determining the severity and duration of symptoms of infection. Polymorphisms in TLR1 and -2 have been linked to altered innate immune responses to *B. burgdorferi*, providing strong evidence that host factors contribute to clearance of the pathogen and modulation of innate defenses, both early and late in infection (7–9). However, these polymorphisms do not explain the extent of disease manifestation in infected patients. The strong familial association of other types of arthritis is consistent with the concept that genetic predisposition may trigger a pathologic response to inflammatory stimuli, such as bacterial infection. Furthermore, certain of the MHC alleles associated with rheumatoid arthritis have been identified as contributing to antibiotic-refractory Lyme arthritis by some groups, suggesting the possibility of other common mechanisms in immune-mediated stages of disease (10–12).

In a seminal observation, Barthold and colleagues reported that inbred strains of laboratory mice exhibit consistent differences in arthritis severity following infection with *B. burgdorferi*, and identified the C3H/HeJ or C3H/HeN (C3H) mouse as displaying the most severe disease and C57BL/6 (B6) as displaying less severe disease (13). Key features of arthritis development that are shared between infected C3H mice and patients with severe Lyme arthritis include synovial hyperplasia, inflammatory cell infiltrate, and exuberant edema. The mouse is one of the major mammalian reservoirs for *B. burgdorferi*, serving as host for blood meals of the larval and nymphal stages of *Ixodes sps* vector ticks (14). Studies in the mouse have been critical to our understanding of environmentally regulated changes in gene expression necessary for transition between hosts and the elaborate mechanism of antigenic variation necessary for survival in the immunocompetent host (15).

The imperative health need for understanding the genetic component of human Lyme arthritis prompted us to pursue a forward genetic approach to identify genes responsible for disease severity. Barthold’s initial studies (13) identified C3H and B6 as being at the extreme ends of Lyme disease symptoms, and more recent SNP-based analysis has placed B6 and C3H mice on well-separated branches of the mouse family tree (16). B6 is the reference strain for the mouse genome sequence, and C3H mice were included in the Perlegen Sciences resequencing project, permitting the extensive analysis of variation between these strains. Our work is the outcome of a classic forward genetic approach to identifying regulators of Lyme arthritis severity in mice. Intercross populations between C3H and B6 (or BALB/c) mice led to the identification of multiple *B. burgdorferi* arthritis-associated (*Bhaa*) quantitative trait loci (QTL) on 5 chromosomes (17). Four individual crosses identified *Bhaa2* on mouse chromosome 5, which exhibits the strongest linkage to disease severity, with a maximum lod score of 10.2 (18). Our laboratory previously developed a B6.C3H-*Bhaa2Bhaa3* congenic mouse line, where *Bhaa2Bhaa3* from susceptible C3H mice was isolated on an otherwise uniform resistant B6 genetic background (19, 20), and found that these mice exhibit increased Lyme arthritis severity, with joint inflammation and histopathology closely resembling the human disease.

Conflict of interest: The authors have declared that no conflict of interest exists.

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Figure 1
Positional mapping and characterization of the \textit{Gusb}+ allele. (A) Advanced congenic lines identify regulatory subintervals within \textit{Bbaa2}. Each row represents the genetic makeup of 1 congenic mouse line across the \textit{Bbaa2} interval (120.3 to 141.2 Mb) on mouse chromosome 5. White and black portions of each row represent areas inherited from the B6 or C3H background, respectively. Ankle swelling measurements taken 4 weeks after \textit{B. burgdorferi} infection (\(n = 12\) to 35 mice per group; overall \(P < 0.0001\)). Significance of cosegregation (right) between ankle swelling and blinded scores of joint histopathology and PMN infiltration, assessed by 1-tailed Mann-Whitney test. (B) Inheritance of the \textit{Gusb} polymorphism among strains included in the Sanger SNP resequencing database. (C) C3H mice and congenics carrying the \textit{Gusb}+ allele exhibited enzymatic hypomorphism in serum and bone marrow-derived macrophage cell extracts and supernatants (\(n = 4\)). (D) CBA/Ca expressed near normal serum GUSB activity, while CBA/J shared the C3H GUSB hypomorphism. (E) CBA/J developed severe Lyme arthritis, while CBA/Ca were resistant (\(n = 5\) [B6 and C3H] and 10 [CBA substrains] mice in each group; overall \(P < 0.0001\)). Significance assessed by 1-way ANOVA followed by Dunnet’s multiple comparison test versus B6 (A and E) or Bonferroni’s post test (C and D). * \(P < 0.05\); ** \(P < 0.01\); *** \(P < 0.001\); **** \(P < 0.0001\).
In this study, we report the positional cloning of a key genetic regulator underlying the increased Lyme arthritis severity conferred by Bbaa2 in C3H mice, the lysosomal enzyme β-glucuronidase, Gusb. The hypomorphic C3H allele, Gusb<sup>h</sup>, was found to cause increased arthritis severity in mouse models of both Lyme and rheumatoid arthritis. Gusb belongs to a recently recognized group of lysosomal enzymes that modulate lysosomal storage and function and that are coregulated in response to stress. We propose that mild deficiencies in Gusb and other coregulated lysosomal enzymes may have previously unrecognized impact on a variety of inflammatory pathologies.

**Results**

**Positional cloning of Gusb.** Through additional backcrossing to the parental B6 line, we developed 15 advanced B6.C3H-Bbaa2 congenic mouse lines harboring subintervals of Bbaa2<sup>C3H</sup> from 120.3 to 141.2 Mbp (Figure 1A and ref. 21). After infection with B. burgdorferi, the various subinterval congenic lines exhibited a wide spectrum of disease severity, as assessed quantitatively by ankle swelling measurements. Compared with B6, congenic mice harboring C3H-derived intervals from 129.0–130.5 Mbp (<i>P</i> < 0.01), 133.5–141.2 Mbp (<i>P</i> < 0.05), and 125.3–128.2 Mbp (<i>P</i> < 0.05) within Bbaa2 exhibited significantly more severe disease. The ankle swelling data also support the presence of a negative regulatory element within Bbaa2 (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI72339DS1). Moreover, for the 129.0–130.5 Mbp and 133.5–141.2 Mbp intervals, increases in the categorical traits of pathology score and neutrophil (PMN) infiltration cosegregated with ankle swelling. Consequently, they have been designated Bbaa2<sup>a</sup> and Bbaa2<sup>b</sup>, respectively.

**Figure 2**

Loss of GUSB function exacerbates Lyme arthritis severity in a genetically recessive manner. (A) Gusb<sup>null</sup> mice do not exhibit a defect in host defense. The observed difference between B6 and C3H genetic backgrounds in heart bacterial burden has been previously described (59). (B) Serum GUSB activity of infected B6 and C3H controls, Gusb<sup>null</sup> homozygotes, and Gusb<sup>heterozygous</sup> littersates (n = 5 to 6 per group). (C) Arthritis severity measurements of Gusb<sup>null</sup> homozygotes, heterozygous littersates, and WT B6 and C3H controls (n = 5 to 6 per group; overall <i>P</i> < 0.0001). (D and E) Arthritis severity measurements of B6.C3H-Bbaa2 and B6.C3H-Gusb<sup>heterozygous</sup> littersates were statistically indistinguishable from B6 control animals (n = 5 per group; overall <i>P</i> < 0.001). Significance assessed by 1-way ANOVA followed by Bonferroni’s multiple comparison test versus B6. *<i>P</i> < 0.05; ***<i>P</i> < 0.001; ****<i>P</i> < 0.0001.
GUSB hypomorphism influences arthritis severity through a cell-intrinsic mechanism. Radiation chimeras were generated between B6 and B6.C3H-Bbaa2 in all pairwise combinations. We achieved high level (>90%) engraftment of B cells and myeloid lineages (Supplemental Figure 6). (A) Chimera serum GUSB activity levels were determined by the donor cell source. (B and C) The C3H Bbaa2 locus contributed to more severe Lyme arthritis, primarily through the activity of radiation resistant joint resident cells. Notably, the B6→Bbaa2 group developed severe Lyme arthritis despite high serum GUSB levels, and the Bbaa2→B6 group was resistant despite low serum GUSB levels (n = 16 to 20 rear ankle joints, 8 to 10 mice per group; overall P < 0.0001). Significance of ankle swelling assessed by 1-way ANOVA followed by Dunnet’s multiple comparison test versus the B6→B6 transplant control. Significance of overall lesion scores assessed by Mann-Whitney test versus the B6→B6 transplant control, with Bonferroni correction. *P < 0.05; **P < 0.01; ***P < 0.0001.

C3H mice carry a natural variant of Gusb that is functionally hypomorphic. The C3H strain is known to carry a functionally hypomorphic Gusb<sup>b</sup> allele, which confers a 70%-90% reduction in enzymatic activity in the serum and various tissues (23, 24). We verified that our B6.C3H-Bbaa2 congenic mice exhibited hypomorphic GUSB enzymatic activity in serum and in cell extracts and supernatants obtained from strain-specific bone marrow-derived macrophages (Figure 1C). We observed no significant differences in Gusb mRNA levels between strains or following <i>B. burgdorferi</i> infection, indicating that this hypomorphism manifests posttranscriptionally (Supplemental Figure 3). The Sanger SNP database indicates that only 3 of the 18 included strains, C3H/HeJ, AKR/J, and CBA/J, share this Gusb coding variant. Both C3H and AKR/J have previously been shown to develop severe Lyme arthritis (13, 25). Intriguingly, the lower density CDG-MDA1 database shows that unlike CBA/J, a closely related CBA/Ca strain carries the common B6 reference nucleotide, which we verified by PCR SNP genotyping (Supplemental Figure 4 and ref. 26). These CBA substrains arose from a partially inbred line and have been genetically isolated but never outcrossed (27), suggesting the Gusb allelic difference is likely to be the result of a limited amount of residual heterozygosity that existed prior to separation. Consistent with this genetic difference, analysis of serum GUSB enzymatic activity levels confirmed that, although CBA/Ca mice had levels similar to those of B6, CBA/J mice were GUSB hypomorphs like C3H mice (Figure 1D). Following <i>B. burgdorferi</i> infection, CBA/J mice developed severe arthritis, while CBA/Ca were resistant (Figure 1E). Importantly, the middensity CDG-MDA1 database identifies only 118 coding nonsynonymous SNPs in the entire genome distinguishing between CBA/Ca and CBA/J mice, and only this Gusb polymorphism was positioned within any of the previously identified Bbaa QTL (17, 19, 28), including the Bbaa2b region referenced in Figure 1A.

Loss of GUSB function exacerbates Lyme arthritis severity in a genetically recessive manner. The availability of a spontaneous Gusb mutant mouse line on the resistant B6 genetic background, B6.C3H-GusbNull<sup>-mps-2J</sup>/BrkJ (GusbNull), allowed us to determine the impact of GUSB loss of function in a second, independent mouse line. This GusbNull strain is also a congenic line (Methods), and GusbNull homozygotes are used as a mouse model of mucopolysaccharidoses type VII (MPSVII). Importantly, we determined that GusbNull mice exhibit no defect in host defense (Figure 2A) despite expressing only 1% of normal GUSB levels in homozygotes (Figure 2B). Infected homozygous GusbNull mice developed maximally severe Lyme arthritis, while heterozygous littermates carrying 1 functional Gusb<sup>b</sup> allele were protected (Figure 2C). We corroborated the finding that heterozygous animals were protected with both our full-length B6.C3H-Bbaa2 and our B6.C3H-Gusb<sup>b</sup> congenic mouse lines (Figure 2, D and E). Thus, neither the severe GusbNull allele nor the hypomorphic Gusb<sup>b</sup> allele act in a dominant negative fashion to interfere with the protective activity of functional Gusb<sup>b</sup> alleles present in these heterozygous animals.

GUSB hypomorphism acts through a cell-intrinsic mechanism. GUSB functions as a lysosomal hydrolase that requires the low pH of the lysosome for full enzymatic activity, but is also present in serum. To clarify which pool of GUSB is responsible for its effect on arthritis, radiation chimeras were generated in all pairwise combinations between B6 and B6.C3H-Bbaa2 congenic mice (29). Although the donor cell source, not the recipient genotype, was identified as responsible for serum GUSB levels in chimeric mice (Figure 3A), Lyme arthritis severity was exacerbated in both the B6→B6.C3H-Bbaa2 and the B6.C3H-Bbaa2→B6.C3H-Bbaa2 groups relative to the B6→B6 control group (Figure 3, B and C).
Notably, the disease severity of the B6→B6.C3H-Bbaa2 group was increased despite high GUSB activity levels in the serum. Conversely, B6.C3H-Bbaa2→B6 chimeric mice did not develop significantly more severe disease than the B6→B6 control group, despite low serum GUSB activity. These results indicate that GUSB hypomorphism primarily modulates disease severity within joint-resident, radiation-resistant cells and that serum GUSB levels are not determinative. This suggests that GUSB hypomorphism acts through a localized, cell-intrinsic mechanism to initiate the development of inflammatory arthritis.

Transgenic overexpression of Gusβ in C3H mice reduces Lyme arthritis severity. Because Gusβ does not appear to interfere with the function of Gusα in a dominant negative fashion in our various heterozygous experiments and because our radiation chimera experiments implicate joint resident cell types in arthritis development, transgenic overexpression to correct GUSB levels in a hypomorphic strain was considered a reasonable approach. To determine the magnitude of the Gusβ effect, a transgene driving ubiquitous mouse Gusβ expression (Figure 4A) was used to produce C3H/HeN-CAG-Gusβ transgenic mice (GusβTg). Five founders were identified that met or exceeded the serum Gusβ enzymatic activity levels of B6 mice (Figure 4B). These GusβTg founders were bred to C3H/HeN mice, and progeny with elevated serum Gusβ enzymatic activity levels were selected for infection (Supplemental Figure 5). Although serum Gusβ enzymatic activity levels were not expected to directly influence disease severity, transgenic progeny were screened for serum Gusβ activity as a facile surrogate metric of the approximate expression level achieved ubiquitously in all tissues. Following infection with B. burgdorferi, we found that GusβTg progeny exhibited a profound and highly significant (P < 0.001) reduction in disease severity (Figure 4C) relative to WT C3H control mice. This argues that among the many Bbaa loci previously identified to regulate Lyme arthritis severity in C3H mice, Gusβ is a key regulator.

Evidence of a conserved role for Gusβ in a model of rheumatoid arthritis. Because the B6.C3H-GusβTg congenic line provides the greatest genetic stringency to interrogate the specific impact of GUSB hypomorphism on a resistant genetic background, we used it to determine whether alterations in Gusβ modulate disease severity in a way that is unique to Lyme arthritis or whether it plays a more generalized role. To test this, we used the K/BxN serum transfer model of rheumatoid arthritis as a second experimental approach to inducing disease (30, 31). This model isolates the downstream effector phase of disease pathogenesis from the initiation phase through adoptive transfer of arthritogenic autoantibodies to induce a joint-specific inflammatory arthritis. Injection of submaximal doses of K/BxN serum was useful in determining the unique contribution of Gusβ to arthritis severity in this model. Following intraperitoneal injections of 100 μl K/BxN serum on days 0 and 2, we found that our B6.C3H-GusβTg congenic mice began to exhibit more severe ankle swelling than B6 control animals beginning on day 4, which was further exacerbated on day 7 (Figure 5A). Histopathology scores for joints at day 7 also corroborated the significance (P < 0.05) of this effect (Figure 5B).

GUSB deficiency is associated with excessive accumulation of glycosaminoglycans during arthritis development. GUSB is a lysosomal hydrolase that catalyzes an essential step in the homeostatic degradation of glycosaminoglycans (GAGs). Severe autosomal recessive GUSB deficiency causes a lysosomal storage disease known as MPSVII, one characteristic of which is spontaneous accumulation of partially degraded GAGs within lysosomes (32). C3H mice begin to develop mild lysosomal accumulation of GAGs by 12 months of age, but younger 9- to 11-week-old mice appear to be unaffected (33). GAGs and partially degraded fragments have previously been implicated as direct mediators of inflammation through activation of TLRs (34). Rodent models of lysosomal storage disease have been shown to exhibit less severe symptoms following the genetic removal of TLR4 or the pharmacological blockade of TNF-α signaling, consistent with an inflammatory component to disease pathogenesis (35, 36). To determine whether GAG accumulation occurs during arthritis development in our GUSB-deficient strains, we performed Alcian blue staining to detect the presence of acidic polysaccharides, including GAGs, in joint histopathology sections at 4 weeks after infection. The inflamed tissues from Gusβnull and GusβTg strains consistently stained intensely positive for the presence of GAGs. Alcian blue-positive material was identified in the periarticular soft tissue, particularly the tendon sheath, of the joints of Gusβnull and GusβTg strains (Figure 6, B, D, F, and H). GAG accumulation was associated with severe tendonitis manifested by acute inflammation composed of dense neutrophilic infiltrates, tendon hyperplasia, and synovial hypertrophy. In contrast, no significant accumulation of Alcian blue–positive material or inflammation was detected in joints from C3H GusβTg and other GUSB-sufficient strains (Figure 6, A, C, E, and G). Similarly, joints from day 7 K/BxN-treated B6 control animals lacked GAG accumulation despite...
It is well established that mutations causing frank deficiencies in GUSB result in spontaneous and overt lysosomal storage disease, specifically demonstrating a key role for a naturally occurring hypomorphic C3H Gusbb allele in Lyme arthritis severity using a scientifically rigorous QTL mapping approach. All tested inbred mouse strains that carry Gusbb are genetically susceptible to severe Lyme arthritis, and introgression of either Gusbb or the more severe Gusbb<sup>homozygote</sup> onto a resistant B6 genetic background confers susceptibility. Importantly, Gusbb<sup>homozygote</sup> homozygous animals develop a maximal Lyme arthritis response, equivalent to that of C3H-positive control animals, while strains with the milder hypomorphism conferred by Gusbb develop significant (<i>P < 0.01</i>) arthritis of intermediate severity. Additionally, correction of the Gusbb deficiency through transgenic overexpression of Gusbb confers significant (<i>P < 0.001</i>) protection to genetically susceptible C3H mice, conclusively and specifically demonstrating a key role for Gusbb.

It is well established that mutations causing frank deficiencies in GUSB result in spontaneous and overt lysosomal storage disease, although mild deficiencies may go unrecognized until late in life and are often misdiagnosed as an inflammatory joint disease (37). Human Gusbb is known to be polymorphic, with over 750 SNPs recorded in the dbSNP database. Forty-nine mutations causing overt disease have been identified in MPSVII patient populations (32), while the prevalence and impact, if any, of other variants remain undefined. Human GUSB enzymatic activity levels in the general population have been shown to exhibit a wide distribution and vary in tissue and serum samples by up to 30-fold (38, 39), differing larger than those observed between high-expressing (B6, CBA/Ca) and low-expressing (C3H, CBA/J) inbred mouse strains used in this study (Figure 1D and ref. 33). GUSB may be uniquely sensitive to mild deficiencies in enzymatic activity, since it has been suggested as the rate-limiting enzyme in the dermatan sulfate degradative pathway (40). Severe deficiencies in individual lysosomal enzymes causing overt disease are rare, with a combined incidence estimated at up to 1 in 5,000 births (41), far below the practical limit of detection of genome wide association studies (GWAS) (minor allele frequency > 0.05) (42). Although human genetic susceptibility to Lyme arthritis has not been investigated by GWAS, it is noteworthy that thus far only a fraction of the genetic variance underlying rheumatoid arthritis has been identified by GWAS (43), accentuating the added value of our QTL mapping approach.

Our studies have identified a naturally occurring, mild subclinical GUSB deficiency that transforms the normally protective local response to <i>B. burgdorferi</i> into a fulminating inflammatory arthritis. Recent literature has brought attention to the persistence of bacterial antigen in host tissues, even following antibiotic regimens that effectively cleared cultivable bacteria (44). Importantly, the increased disease severity we have observed occurs in the absence of significant alterations in host defense or <i>B. burgdorferi</i> load in tissues (Figure 2A). This disease exacerbation is also retained in the distinct and well-characterized K/BxN serum transfer model of rheumatoid arthritis, where dosage of the inflammatory stimulus can be tightly controlled. This suggests that inflammatory initiators such as <i>B. burgdorferi</i> antigen or autoantibodies trigger severe disease through a 2-hit phenomenon, where coincident breakdown of host tolerance mechanisms designed to limit the pathological consequences of infection and the ensuing inflammatory response instead exacerbate disease symptoms. The ability of Gusbb to exacerbate arthritis in response to very different experimental stimuli suggests this phenomenon may also be generalizable to other inflammatory triggers.

Our observation that radiation-resistant, joint-resident cells are a primary determinant of the magnitude of the Lyme arthritis response offers important insight into the mechanisms underlying joint pathogenesis in this model. This finding indicates that resident cells have important roles both in recruiting inflammatory immune cells to help clear infection and in mitigating damage through tolerance mechanisms. Our complementary finding that the severe joint pathology observed in infected B6.C3H-Bhaa2 mice is not effectively corrected by high-serum GUSB levels bears striking resemblance to reports on the limited efficacy of enzyme replacement therapy to alleviate musculoskeletal symptoms in adult animal models of MPSVII (45, 46), although early intervention in neonates has shown promise (47, 48). Similarly, the joint pathologies in patients with a variety of mucopolysaccharidoses are difficult to treat and respond...
much more slowly to high-dose enzyme replacement therapy than other symptoms, such as hepatosplenomegaly or sleep apnea (49). Taken together, these findings highlight the importance of the primary response to bacterial stimulation that is mounted by resident cells in these refractory joint tissues.

The evident link between GUSB hypomorphism and excessive deposition of GAGs with potential proinflammatory activity provides a plausible mechanism bridging disease to the critical catalytic role GUSB plays in homeostatic GAG degradation. Although our data show no significant change in bacterial load due to GUSB deficiency, GAG-mediated cell adhesion by *B. burgdorferi* does play a noteworthy role in mammalian infection and tissue localization (50). The GUSB substrate dermatan sulfate has been linked to excessive TNF-α release by chondrocytes (40), and MPSVI symptoms are alleviated by blocking TNF-α (36), a highly successful target for rheumatoid arthritis (51). However, this does not preclude the involvement of other downstream effectors. The release and accumulation of lysosomal exoglycosidases in the serum has been observed in multiple forms of chronic inflammatory arthritis, with localized release into synovial fluid reported to be especially exaggerated in chronic Lyme arthritis patients (52, 53). Although lysosomal exoglycosidases such as GUSB are catalytically inactive at neutral pH, coincident release of other proinflammatory lysosomal components may provide an alternate mechanism to trigger or amplify a local inflammatory cascade (54). We suggest that the identification of Gusb as a key regulator of murine Lyme and rheumatoid arthritis severity provides a sound scientific basis for future investigations into serum GUSB or GAG levels as potential biomarkers of human susceptibility to developing chronic or severe inflammatory arthritis.

Gusb is 1 member of a large group of over 40 coregulated lysosomal enzymes in the coordinated lysosomal expression and regulation (CLEAR) network that are responsible for the stepwise degradation of several distinct biological substrates, including GAGs, lipids, sugars, chitin, and glycogen (55). As with Gusb, severe deficiencies in virtually all of these enzymes induce spontaneous lysosomal storage disease. Many such lysosomal storage diseases exhibit progressive joint disease. Recent work has demonstrated that overexpression of the master regulatory transcription factor of the CLEAR network, TFEB, leads to successful clearance of glycogen from lysosomes in both in vitro and mouse models of Pompe disease (56). Based on the close regulatory and functional interrelationship between *Gusb* and other lysosomal enzymes, we propose that the increased arthritis severity observed in this study may also be generalizable to mild deficiencies in other members of the CLEAR network.

**Figure 6**
GUSB enzymatic hypomorphism is associated with exaggerated accumulation of GAGs in the inflamed joint. Representative images of Alcian blue-stained rear ankle joint sections from *B. burgdorferi*-infected GUSB-sufficient (upper panels: A, C, E, and G) or GUSB-deficient (lower panels: B, D, F, and H) strains, or day 7 K/BxN-treated B6 (I) and B6.C3H-Gusb+ (J) congenic mice. Arrowheads indicate position of the cranial tibial tendon sheath. Original magnification, ×4. Scale bars: 500 μm (K) GAG accumulation in the soft tissue and joint/synovial space was scored on a scale of 0–4 (n = 3 to 4 joints per group). Pairwise significance assessed by 1-tailed Mann-Whitney test. *P < 0.05.
Methods

Generation of interval-specific congenic lines
Interval-specific congenic lines (ISCL) were generated as described (19). Standard nomenclature for the mouse lines is used herein, as follows: the background strain (e.g., C57BL/6NCr, abbreviated B6 in the ISCL designations) is listed first, followed by the donor strain (e.g., C3H) and the introgressed interval on mouse chromosome 5 (Mb, according to the NCBI37/38 Mouse Genome Assembly) from C3H/HeN into the C57BL/6NCr parental strain (National Cancer Institute). B6.C3H-120.3-131.0, B6.C3H-125.3-131.8, B6.C3H-129.0-130.5 (B6.C3H-Gusb+), B6.C3H-129.0-141.2, B6.C3H-131.8-133.5, B6.C3H-131.8-141.2, and B6.C3H-134.7-141.2 ISCL were generated by marker-assisted selection using high-resolution melting analysis SNP genotyping primers as described (21). Homozygous progeny derived from matings between heterozygous male and female ISCL mice that were free of background donor strain contamination were used to fix the lines.

GusbNull mice
GusbNull mice were obtained from Jackson Laboratory, as a mouse model of MPSVII. GusbNull mice were derived from heterozygous breeder pairs, and homozygous offspring were identified by markedly sub-Mendelian ratios (data not shown) by SNP genotyping and reduced body size. As per Jackson Laboratory documentation, this spontaneous mutant mouse line originally arose in the C3H/HeOuJ stock production colony and was extensively backcrossed to C57BL/6J. As a result, these mice are congenic for the C3H/HeN-CAG-Gusbb transgenic mice.

Generation of C3H/HeN-CAG-Gusbb transgenic mice
All PCR steps were performed for 25 cycles with high-fidelity Phusion PCR amplified with an annealing temperature of 71°C. The following sequence-specific primers containing 5′ EcoRI cloning sites progressively amplified a 2.2-kb fragment of the predicted size, which was then separated from the backbone by agarose gel electrophoresis. The backbone was excised and gel purified. The purified Gusb+ ORF fragment was cloned into pCAGGS EcoRI and validated by fully sequencing across the insert.

Gusb transgenic mice were obtained from Jackson Laboratory, as a mouse model of MPSVII. Gusb mice were derived from heterozygous breeder pairs, and homozygous offspring were identified by markedly sub-Mendelian ratios (data not shown) by SNP genotyping and reduced body size. As per Jackson Laboratory documentation, this spontaneous mutant mouse line originally arose in the C3H/HeOuJ stock production colony and was extensively backcrossed to C57BL/6J. As a result, these mice are congenic for the C3H/HeN-CAG-Gusbb transgenic mice.

Gusb enzymatic activity assay
4-Methylumbelliferyl β-D-glucuronide (MUG) (Marker Gene) was used as a fluorogenic substrate to measure GUSB enzymatic activity. 10 μl of sample (serum, cell extract, or supernatant) was incubated with 1 μM MUG in a total volume of 50 μl assay buffer (200 mM sodium acetate; pH 4.5; 10 mM EDTA; 0.01% BSA; 0.1% Triton X-100) for 1 hour at 37°C in a 96-well plate (#3370; Costar). 150 μl of stop buffer (200 mM sodium carbonate) was then added, and samples were analyzed on a Biotek Synergy HT microplate reader. Fluorescence was measured with an excitation wavelength of 360 nm and an emission wavelength of 460 nm. Units were calculated by comparison against a standard curve prepared using purified bovine liver glucuronidase (type B-1 #G0251; Sigma-Aldrich). Measurements of β-galactosidase activity were performed in a similar manner using 4-methylumbelliferyl β-D-galactopyranoside (Marker Gene) as a substrate.

Culture and analysis of bone marrow–derived macrophages
BMMφ were prepared as described (19) by culture in RPMI 1640 medium (Invitrogen) supplemented with 30% L929 conditioned medium and 20% horse serum (HyClone). Harvested macrophages were replated into 12-well plates at a density of 6 × 105/ml in medium lacking serum and containing 1% Nutridoma. Supernatants were harvested 24 hours later, cells were washed once with 1× PBS, and cell extracts were then harvested by incubation in extraction buffer (50 mM NaPO4, pH 7.0; 10 mM BME; 10 mM EDTA; 0.1% sarcosyl; 0.1% Triton X-100).

Culture of B. burgdorferi and infection of mice
Mice between 6 and 7 weeks of age were infected by intradermal injection with 2 × 107 bacteria of the B. burgdorferi N40 isolate (provided by Stephen Barthold, UCD, Davis, California, USA). B. burgdorferi cells were cultured in Barbour-Stoenner-Kelly II medium containing 6% rabbit serum (Sigma-Aldrich).

Arthritis analysis
Rear ankle joints were measured at the time of infection and at 4 weeks after infection by using a metric caliper, as described (19). Measurements of the thickest anteroposterior portion of the ankle with the joint extended were taken and are reported as the change in ankle swell-
The Journal of Clinical Investigation

How many people get Lyme disease?

Lyme disease affects an estimated 300,000 people annually in the United States. The disease is caused by the bacterium Borrelia burgdorferi, which is transmitted to humans through the bite of an infected deer tick. Lyme disease can be prevent by taking preventive measures, such as avoiding tick-infested areas and wearing protective clothing.

The disease is characterized by a bulls-eye rash, joint pain, and fatigue. Early diagnosis and treatment with antibiotics can prevent long-term complications. The majority of cases are diagnosed in the northeastern United States, with the highest incidence rates in areas with large populations of deer and blacklegged ticks.

The Journal of Clinical Investigation

Generation and analysis of radiation chimeras

Chimeras were generated in all pairwise combinations between B6 CD45.1 and B6.C3H-Bhsaa2 (CD45.2) congenic mice as described (29). Briefly, 4-week-old mice were lethally irradiated with 2 doses of 525 cGy given 3 hours apart using a GE Isovolt Titan (GE Healthcare). 24 hours later, splenocytes were prepared from donor mice. Irradiated mice each received an intravenous injection of 2 × 10⁶ splenocytes in a 200 μl volume. Chimerism was evaluated at 3 weeks after transplant by flow cytometric analysis of blood leukocytes (Supplemental Figure 6).

K/BxN serum transfer

K/BxN serum was a gift from Paul Allen (Washington University). Rear ankle joints were measured with a metric caliper prior to treatment, as described above. 100 μl of K/BxN serum was administered by intraperitoneal injections on days 0 and 2. Ankle swelling was determined by measurements on days 1, 2, 4, and 7. After the final day 7 measurement, joint histopathology was assessed, as described above.

Imaging of joint histology sections

Alican blue-stained sections were visualized on an Olympus BX41 clinical microscope (Olympus America) using x4 total magnification. Images were recorded with an Olympus DP72 camera and prepared using Olympus CellSens digital imaging software. Images were analyzed for cell infiltration, tendon sheath thickness, and reactive-reparative responses. To assess GAG accumulation, joint sections from each group were given a subjective score based on the presence of Ailcan blue-positive material in the soft tissue and joint/synovial space, ranging from 0 (none) to 4 (severe).

Statistics

All data represent mean ± SEM. All statistical calculations were performed using GraphPad Prism 5. P < 0.05 was considered statistically significant. Continuous variables were analyzed by 1-way ANOVA and Student’s t test. Categorical variables were analyzed by Kruskal-Wallis or Mann-Whitney nonparametric tests. All calculations of Student’s t tests and Mann-Whitney tests are 2-tailed unless otherwise specified.

Study approval

All study protocols involving mice were conducted in accordance with the NIH guidelines for the care and use of animals and approved by the Institutional Animal Care and Use Committee at the University of Utah.

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The Journal of Clinical Investigation

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319


