Higher eukaryotes are colonized by microbial organisms, predominantly on interfaces with the environment such as the skin and gastrointestinal tract (1). Recent studies suggest that host cells, including professional antigen-presenting cells, “sense” the presence of microbial molecules under homeostatic conditions (2, 3). This sensing is performed by several families of cellular receptors, including Toll-like receptors (TLRs) and non-TLR proteins such as NOD/CATERPILLAR family proteins (4, 5). Collectively, these proteins are thought to bind to conserved pathogen-associated molecular patterns (PAMPs) and activate proinflammatory signals in host cells. The appreciation that microbial sensors such as TLRs may chronically or tonically engage PAMPs during homeostasis raises a major question: if TLRs trigger both adaptive and innate immune responses, why do homeostatic TLR signals not trigger inflammation?

Despite the presence of up to $10^{11}$ bacteria per gram of luminal contents within the gastrointestinal tract, most humans tolerate this colonization without adverse effects. Indeed, host cell sensing of luminal microbes may provide beneficial signals under homeostatic conditions (2). How immune responses to commensal flora are restrained during homeostasis thus remains one of the central questions in immune regulation. Inflammatory TLR signals might be distinguished from homeostatic TLR signals by the quantity or strength of TLR signals, by cell type–specific responses, by the cellular location of PAMP/TLR interactions (e.g., apical vs. basal surfaces or intracellular vs. membrane-bound receptors), or by additional extracellular cues (e.g., cellular stress response proteins) that modify the cellular interpretation of such signals. Among these general mechanisms, perhaps the most attention has been focused on proteins that regulate the intracellular strength, duration, and/or character of PAMP-triggered signaling.

TLR proteins contain leucine-rich repeat regions that mediate ligand binding. TLRs also
contain intracellular Toll/IL-1 receptor (TIR) domains that
triger signals via homotypic interactions with proxi-Mal
proteins (6, 7). All TLRs signal via the adaptor MyD88,
leading to activation of NF-κB and mitogen-activated protein
(MAP) kinase pathways. NF-κB activation leads to a tran-
scriptional program that up-regulates proinflammatory gene
products such as IL-1β and TNF-α and, ultimately, to the ac-
tivation and accumulation of monocytes. MyD88 is essential
for most TLR-induced NF-κB signaling, as MyD88-deficient
(MyD88−/−) cells exhibit reduced and delayed NF-κB signaling
activity (8, 9). TLR3 and TLR4 also use another adaptor,
TIR domain–containing adaptor inducing IFN-β (TRIF).
TRIF-dependent signals activate IFN response factor 3 (IRF3)
in addition to NF-κB (10, 11). These transcription factors
induce productive antiviral immune responses by binding to
promoters of type I IFN and chemotactic genes (12).

Constitutive MyD88-dependent TLR signals provide
beneficial, noninflammatory signals (2). These signals can also
triguer spontaneous inflammation. Thus, a major issue arises
as to how homeostatic TLR signals are distinguished from in-
flammatory TLR signals. One potential mechanism for distin-
guishing these TLR signals may be related to recent discoveries
that endogenous proteins such as IL-1R–associated kinase M
(IRAK-M), ST2, and single Ig and TIR domain (SIGIRR)
can restrict TLR signals (13–18). These proteins can restrict the
duration and/or intensity of TLR signals and modulate the
cellular outcome of TLR signaling, thereby helping to
determine whether TLR signals lead to homeostatic or in-
flammatory responses. Notably, mice lacking IRAK-M, ST2,
and SIGIRR exhibit relatively modest inflammation and
survive for longer than 9–12 mo (14, 16, 17). Suppressor
of cytokine signaling (SOCS) 1 can also inhibit TLR signaling,
and SOCS-1−/− mice exhibit severe spontaneous inflamm-
ation and lymphocyte-mediated perinatal lethality; however
SOCS-1−/− mice are rescued by the absence of IFN-γ signaling
in SOCS-1−/− IFN-γ−/− double-mutant mice (18, 19).
Thus, it is unclear whether these proteins restrict TLR signals
in unperturbed mice and whether restricting homeostatic
TLR signals is important for immune homeostasis.

A20 is an inducible ubiquitin-editing enzyme that restricts
both TLR- and TNF-induced responses by regulating the
ubiquitylation of key signaling proteins (20, 21, 22, 23). A20-
deficient (A20−/−) mice spontaneously develop multiorgan
inflammation, severe cachexia, and premature lethality that
can start within 1–2 wk of age, demonstrating the potent
antinflammatory functions of this molecule (22). In addition,
A20−−/− RAG-1−/− double-mutant mice spontaneously de-
velop severe inflammation, cachexia, and premature death,
indicating that A20 potently regulates innate immune homeo-
Stasis in the absence of adaptive lymphocytes (22). Although
A20 is critical for restricting TNF-induced NF-κB signals,
both A20−/− TNF−/− and A20−/− TNFR1−/− double-mutant
mice spontaneously develop inflammation, cachexia, and
premature death, indicating that A20 is critical for restrict-
ing inflammatory signals independently of TNF/TNFR1 signal-
ing (22).

A20 expression is rapidly induced in macrophages upon
TLR stimulation and is required for restricting the duration
of TLR–induced NF-κB signaling and the quantity of TLR-
induced proinflammatory cytokines (22). Hence, A20 functions
as a negative feedback regulator of exogenous TLR–induced
signals. As TLR ligands from commensal microorganisms may
engage host cells during homeostatic conditions, we have
investigated the potential roles A20 may play in regulating
homeostatic TLR signals by interbreeding A20−/− mice with
MyD88−/− mice. In addition, we have questioned the role of
commensal bacteria in driving spontaneous inflammation in
A20−/− mice. Finally, we have used A20−/− MyD88−/− double-
mutant mice to examine the role of A20 in regulating MyD88-
independent TLR signals.

RESULTS

MyD88 deficiency prevents premature lethality
and cachexia in A20−/− mice

As commensal microbes or endogenous ligands may engage
host TLRs under basal conditions, we investigated whether
TLR signals contribute to the development of spontaneous
inflammation in A20−/− mice by genetically removing MyD88-
dependent TLR signals. A20−/− mice were interbred with
MyD88−/− mice and backcrossed at least five generations to
C57BL/6. Remarkably, although only 30% of A20−/− mice
survive to 8 wk of age, nearly 95% of A20−/− MyD88−/− double-
mutant mice survive past this age (Fig. 1 A). This improved
survival rate is similar to that of MyD88−/− mice. Additionally,
surviving A20−/− MyD88−/− double-mutant animals do not
exhibit the severe cachexia manifest by surviving A20−/− mice
(Fig. 1 B). A20−/− mice weigh only ~60% of what wild-
type controls weigh, whereas A20−/− MyD88−/− mice weigh
~90% of what MyD88−/− controls weigh (Fig. 1 C). These
data suggest that MyD88 signals drive potent proinflammatory
responses that lead to cachexia and premature death in the
absence of A20.

Homeostatic MyD88-dependent signals activate myeloid
cells in A20−/− mice

MyD88-dependent TLR signals activate myeloid cells, leading
to the production of cytokines that recruit and activate other
innate immune cells. A20−/−, A20−/− RAG-1−/− double-mutant,
A20−/− TNF−/− double-mutant, and A20−/− TNFR1−/− double-
mutant mice all spontaneously accumulate myeloid cells in
both lymphoid and nonlymphoid tissues (21, 22). To begin
to examine the role of MyD88 in causing inflammation in A20−/−
mice, we measured the spleen weights as a reflection of the cellu-
larity of these organs. Spleen weights were dramatically increased
in A20−/− mice when compared with wild-type or MyD88−/−
mice (Fig. 2 A), and this splenomegaly was particularly impres-
sive given that A20−/− mice were significantly smaller than the
other genotypes of mice (Fig. 1 C). In contrast, the weight of
spleens obtained from double-deficient A20−/− MyD88−/− mice
was significantly less than that obtained from A20−/− mice and
more closely approximated the weight of spleens obtained from
wild-type and MyD88−/− mice (Fig. 2 A).
MyD88-dependent signals. MyD88-dependent signals on antigen-presenting cells support the activation and expansion of T cells. To determine whether basal MyD88-dependent signals cause spontaneous activation and expansion of T cells in unperturbed A20−/− mice, we examined the number and activation state of T cells from A20−/−, A20−/− MyD88−/−, and control mice. Significantly greater percentages (and total numbers) of activated, memory phenotype (CD44hi CD62Llo) T cells were present in lymph nodes and spleens of A20−/− mice when compared with wild-type mice (Fig. 3 A and not depicted). In contrast, T cells from A20−/− MyD88−/− double-mutant mice were less activated than those from A20−/− mice and were more comparable to wild-type and MyD88−/− mice. This reduction in spontaneous T cell activation was observed in both CD4+ and CD8+ T cell populations (Fig. 3 B and C). This finding suggests that homeostatic MyD88-dependent signals cause T cell activation in the absence of A20.

**Homeostatic MyD88-dependent signals in radiation sensitive hematopoietic cells drive myeloid cell activation in A20−/− mice**

Hematopoietically derived cells such as macrophages and dendritic cells are able to sample surrounding microflora and transmit basal proinflammatory signals. In addition, A20 and MyD88 are both expressed in epithelial cells and stromal cells, which can also respond to TLR ligands. Thus, A20’s role in restricting MyD88-dependent signals could involve signals on either hematopoietic cells, stromal cells, or both. To isolate the potential role of A20 in regulating MyD88 signaling in hematopoietic cells, Ly5.2 C57BL/6 mice were lethally irradiated and reconstituted with bone marrow hematopoietic stem cells (HSCs) from either wild-type, A20−/−, MyD88+/+, MyD88−/−, or A20−/− MyD88−/− mice. Flow cytomeric analyses of tissues from these chimeras were then performed 6 wk after reconstitution. Similar to our findings with intact mice, spleens from radiation chimera reconstituted with A20−/− HSCs contained increased numbers of various myeloid cell types when compared with chimera reconstituted with wild-type HSCs (Fig. 4 A).
These indicators of spontaneous inflammation were absent in A20−/− MyD88−/− HSC-reconstituted chimera (Fig. 4 B). As IL-10-deficient mice have recently been described to develop spontaneous colitis that resolves in the absence of MyD88 signals (3), we examined whether A20−/− HSC-reconstituted chimera were IL-10 deficient. A20−/− mice expressed elevated levels of IL-10, suggesting that IL-10 deficiency does not contribute to spontaneous inflammation in A20−/− mice (Fig. 4 B). A20−/− HSC-reconstituted chimera also expressed normal levels of IL-2 and possessed normal numbers of functional regulatory T cells (Fig. 4 B and not depicted). Thus, IL-2 deficiency is also unlikely to explain the spontaneous inflammation in A20−/− mice. Finally, A20−/− HSC-reconstituted chimera expressed elevated levels of IL-13 as well as the prototypical Th1 cytokines TNF and IFN-γ (Fig. 4 B and not depicted). Hence, no gross evidence of Th1 or Th2 skewing is evident in these mice.

The myeloid cells that expand in A20−/− HSC-reconstituted chimera spleens include various myeloid populations such as macrophages (Mac-1+, F480+, and Siglec F−), neutrophils (Mac-1+, Gr-1+, F480+, and Siglec F−), and eosinophils (Mac-1+, Siglec F+, and SSCHi) without any particular skewing toward particular subsets (Fig. 4 A). The absolute numbers of these populations were generally reduced in A20−/− MyD88−/− HSC-reconstituted chimera compared with A20−/− HSC-reconstituted chimera, although some of these reductions were less dramatic than the differences observed in intact A20−/− MyD88+/+ and A20−/− MyD88−/− mice (Figs. 2 and 4 A).

To further characterize the nature of the spontaneous inflammation caused by A20−/− hematopoietic cells in these chimera, we analyzed splenic production of cytokines. These studies revealed that spleens from A20−/− HSC-reconstituted chimera produced elevated levels of TNF and IL-1β (Fig. 4 B).
orders of magnitude fewer intestinal bacteria (not depicted). In contrast with A20⁻/⁻/HSC-reconstituted chimera treated with trimethoprim-sulfamethoxazole alone, A20⁻/⁻/HSC-reconstituted chimera treated with the cocktail of broad spectrum antibiotics maintained their weight (Fig. 5 A). Thus, antibiotic treatment prevents cachexia caused by A20 deficiency. Large numbers of Mac1⁺ Gr1⁺ myeloid cells spontaneously accumulated in the spleens of A20⁻/⁻/HSC-reconstituted chimera but not in spleens from antibiotic-treated A20⁻/⁻/HSC-reconstituted chimera (Fig. 5 B). Finally, spleens from A20⁻/⁻/HSC-reconstituted chimera spontaneous produced elevated levels of IL-6 and TNF messenger RNAs (mRNAs) compared with spleens from wild-type HSC-reconstituted chimera (Fig. 5 C). These indicators of inflammation were ameliorated in A20⁻/⁻/HSC-reconstituted chimera treated with broad spectrum antibiotics (Fig. 5 C). Collectively, these results indicate that commensal intestinal bacteria drive spontaneous cachexia and myeloid inflammation via MyD88 signals in the absence of A20.

Collectively, these data suggest that homeostatic MyD88 signals in hematopoietic cells drive a broad spectrum of spontaneous proinflammatory pathways in the absence of A20.

Commensal bacteria stimulate homeostatic MyD88-dependent signals

Homeostatic MyD88-dependent signals may be derived from host molecules or from commensal microbes that express TLR ligands and are in constant contact with host cells. By far, the most abundant source of commensal microbes is the intestinal lumen. Thus, to determine whether commensal intestinal microbes stimulate the homeostatic MyD88-dependent signals that drive inflammation in A20⁻/⁻/ mice, we first generated radiation chimera using HSCs from A20⁺/+ and A20⁻/⁻ bone marrows. 2 wk after irradiation, we treated these chimera with oral broad spectrum antibiotics previously shown to markedly reduce the number of intestinal bacteria (vancomycin, neomycin, metronidazole, ampicillin, and trimethoprim-sulfamethoxazole). After 3 wk of treatment with this antibiotic cocktail (or with trimethoprim-sulfamethoxazole alone as a control), chimeric mice were analyzed for the numbers of intestinal bacteria as well as for evidence of spontaneous inflammation. Chimera reconstituted with A20⁻/⁻ HSCs developed spontaneous cachexia and lost weight when compared with chimera reconstituted with wild-type HSCs (Fig. 5 A). These results suggest that A20⁻/⁻ hematopoietic cells recapitulate the spontaneous inflammation observed in intact A20⁻/⁻/ mice. Chimera treated with the cocktail of broad spectrum antibiotics possessed several

![Figure 3. MyD88 dependence of T cell hyperactivation in A20⁻/⁻/ mice.](image)

In contrast with A20⁻/⁻/HSC-reconstituted chimera treated with trimethoprim-sulfamethoxazole alone, A20⁻/⁻/HSC-reconstituted chimera treated with the cocktail of broad spectrum antibiotics maintained their weight (Fig. 5 A). Thus, antibiotic treatment prevents cachexia caused by A20 deficiency. Large numbers of Mac1⁺ Gr1⁺ myeloid cells spontaneously accumulated in the spleens of A20⁻/⁻/HSC-reconstituted chimera but not in spleens from antibiotic-treated A20⁻/⁻/HSC-reconstituted chimera (Fig. 5 B). Finally, spleens from A20⁻/⁻/HSC-reconstituted chimera spontaneous produced elevated levels of IL-6 and TNF messenger RNAs (mRNAs) compared with spleens from wild-type HSC-reconstituted chimera (Fig. 5 C). These indicators of inflammation were ameliorated in A20⁻/⁻/HSC-reconstituted chimera treated with broad spectrum antibiotics (Fig. 5 C). Collectively, these results indicate that commensal intestinal bacteria drive spontaneous cachexia and myeloid inflammation via MyD88 signals in the absence of A20.

TRIF-dependent TLR signals are restricted by A20

In the absence of the TLR adaptor MyD88, TLR3 and TLR4 signal through an alternate adaptor protein, TRIF. TRIF-dependent responses lead to the production of type I IFNs (IFN-α and IFN-β) and are critical for productive antiviral immunity (10, 11). We thus asked whether A20 was physiologically required for restricting TRIF-dependent TLR responses. To examine this question, we injected LPS intraperitoneally

![Figure 3. MyD88 dependence of T cell hyperactivation in A20⁻/⁻/ mice.](image)
Figure 4. A20 expression in hematopoietic cells restricts MyD88-driven spontaneous inflammation in radiation chimera. (A) Flow cytometric analyses of chimeric mice generated by transfer of A20+/+ MyD88+/+, A20−/− MyD88+/+, A20+/+ MyD88−/−, or A20−/− MyD88−/− bone marrow HSCs into lethally irradiated C57BL/6J mice. 6 wk after reconstitution, spleens from chimeric mice were analyzed by flow cytometry for the number of the indicated myeloid cell types (Mac1/+, Gr1/+, activated macrophages and neutrophils; Mac1+/+ Gr1−/+, neutrophils; Mac1− F4/80+/+ Siglec F−, macrophages; and F4/80− Siglec F+/+ SSCHi, eosinophils). The numbers of Mac−1+, Mac−1+ Gr−1+, and Mac−1+ F4/80− Gr−1+ cells were statistically different (P < 0.05) between A20−/− and A20+/+ MyD88−/− mice. (B) Quantitative real-time PCR analyses of mRNA levels of the indicated cytokines in spleens from chimeric mice. Splenic expression levels of the indicated cytokines are shown in R.U. for the chimeric mice described in A. All cytokine mRNA levels were normalized to β-actin mRNA. The levels of cytokines were statistically different between A20+/+ and A20−/− MyD88−/− mice for all cytokines. Error bars indicate standard deviations. Graphs display results from at least three independent mice per genotype.
Mice were generated from transfer of A20+/+ (WT) or A20−/− bone marrow HSCs into sublethally irradiated congenic Ly5.2+ C57BL/6J mice. 2 wk after irradiation, animals were given 0.5 gram per liter of vancomycin, 1 gram per liter of ampicillin, 1 gram per liter of neomycin, 1 gram per liter of metronidazole, and trimethoprim-sulfamethoxazole in drinking water for 3–4 wk (+Abx) or maintained on trimethoprim-sulfamethoxazole (Ctrl) for the same period. (A) The percent weight change of chimeric mice reconstituted with HSCs of the indicated genotypes and treated with either broad spectrum (+Abx) or control antibiotics (Ctrl). The percent change was calculated by dividing the weight of each mouse on the day of death/analysis by the weight on the day of irradiation and HSC reconstitution. (B) Flow cytometric analyses of Ly5.1+ Mac1+ Gr1+ cells from spleens of chimeric mice. (C) Quantitative real-time PCR analysis of mRNA levels of IL-6 and TNF in spleens of chimeric mice. Note that A20−/− HSC-reconstituted chimera spontaneously lose weight, accumulate myeloid cells, and express higher levels of splenic IL-6 and TNF than A20+/+ HSC-reconstituted chimera, whereas treatment of A20−/− HSC-reconstituted chimera with broad spectrum antibiotics prevents these signs of inflammation. Error bars represent standard deviations. Data are representative of five mice.

### Figure 5. Depletion of commensals with broad-spectrum antibiotics diminishes inflammation driven by A20−/− hematopoietic cells. Chimeric mice were generated from transfer of A20+/+ (WT) or A20−/− Ly5.1+ bone marrow HSCs into sublethally irradiated congenic Ly5.2+ C57BL/6J mice. 2 wk after irradiation, animals were given 0.5 gram per liter of vancomycin, 1 gram per liter of ampicillin, 1 gram per liter of neomycin, 1 gram per liter of metronidazole, and trimethoprim-sulfamethoxazole in drinking water for 3–4 wk (+Abx) or maintained on trimethoprim-sulfamethoxazole (Ctrl) for the same period. (A) The percent weight change of chimeric mice reconstituted with HSCs of the indicated genotypes and treated with either broad spectrum (+Abx) or control antibiotics (Ctrl). The percent change was calculated by dividing the weight of each mouse on the day of death/analysis by the weight on the day of irradiation and HSC reconstitution. (B) Flow cytometric analyses of Ly5.1+ Mac1+ Gr1+ cells from spleens of chimeric mice. (C) Quantitative real-time PCR analysis of mRNA levels of IL-6 and TNF in spleens of chimeric mice. Note that A20−/− HSC-reconstituted chimera spontaneously lose weight, accumulate myeloid cells, and express higher levels of splenic IL-6 and TNF than A20+/+ HSC-reconstituted chimera, whereas treatment of A20−/− HSC-reconstituted chimera with broad spectrum antibiotics prevents these signs of inflammation. Error bars represent standard deviations. Data are representative of five mice.

Finally, IFN-β stimulates IL-15 and IL-15Rα production, leading to NK cell activation in early antiviral immunity. Soluble IL-15Rα is released into the serum during this process (unpublished data). We thus measured serum IL-15Rα protein levels in LPS-stimulated chimeric mice by ELISA, and these assays showed that higher levels of soluble IL-15Rα were present in LPS-stimulated A20−/− MyD88−/− chimera than in LPS-stimulated MyD88−/− chimera (Fig. 6 D). Collectively, these findings indicate that A20 is physiologically required for restricting acute, MyD88-independent, TRIF-dependent proinflammatory signals in vivo.

**A20 regulates TRIF-dependent TLR-induced NF-κB signaling but not IRF3 signaling**

TRIF-dependent TLR signals involve both NF-κB and IRF3-dependent signals (26, 27). To determine how A20 regulates TRIF-dependent TLR responses, we first interbred A20−/− MyD88−/− mice with TNF−/− mice. The resulting triple-mutant mice allowed us to evaluate TRIF-dependent TLR signals in the absence of secondary TNF signaling. We then prepared bone marrow–derived macrophages (BMDMs) from these TNF−/− MyD88−/− and A20−/− TNF−/− MyD88−/− mice.
for the expression of IκBα and phospho-IκBα by immunoblotting. When compared with TNF−/− MyD88−/− BMDMs, A20−/− TNF−/− MyD88−/− cells exhibited prolonged expression of phospho-IκBα and delayed recovery of IκBα, indicating that A20 is required for restricting TRIF-dependent NF-κB signaling (Fig. 7 B).

In addition to activating NF-κB signaling, TLR3- and TLR4-induced activation of TRIF also causes IRF3 phosphorylation, which in turn leads to its dimerization, nuclear translocation, and stimulation of IFN-γ gene transcription (12). TNF−/− MyD88−/− and A20−/− TNF−/− MyD88−/− BMDMs were stimulated with LPS for various time periods. Phospho-IRF3 levels were then measured by immunoblotting of nuclear protein extracts. Surprisingly, TNF−/− MyD88−/− and A20−/− TNF−/− MyD88−/− BMDMs secreted greater amounts of IFN-γ than TNF−/− MyD88−/− BMDMs (Fig. 7 A and not depicted). Thus, A20 appears to be directly required for restricting TRIF-dependent TLR responses independently of TNF signaling.

The production of IFN-γ is dependent on both NF-κB as well as IRF3 signaling, as the IFN-γ promoter contains binding elements for both these transcription factors. To further understand the mechanism by which A20 restricts TRIF-dependent TLR signaling, we stimulated A20−/− TNF−/− MyD88−/− and TNF−/− MyD88−/− BMDMs with LPS, prepared whole-cell lysates at various time points, and analyzed these lysates for the expression of IκBα and phospho-IκBα by immunoblotting.
deubiquitylate TRAF6 (22, 30). Thus, TRAF6 may be a physiological target for A20 during TLR signaling. To directly determine if A20 is essential for regulating TRAF6 ubiquitylation during TLR stimulation, A20+/+ and A20−/− BMDMs were stimulated with LPS for various periods of time, after which cell lysates were analyzed for their expression of ubiquitylated TRAF6. Endogenous TRAF6 is ubiquitylated after 20 min of LPS stimulation (Fig. 8). Importantly, whereas TRAF6 ubiquitylation is transiently induced in A20+/+ cells, declining after 40 min, TRAF6 ubiquitylation is prolonged to 40 and 60 min in A20−/− cells (Fig. 8). As these lysates were boiled in 1% SDS before immunoprecipitation of TRAF6, the ubiquitylated proteins we observe are likely to be predominantly ubiquitylated TRAF6 rather than noncovalently associated proteins. This finding indicates that A20 is essential for restricting endogenous TRAF6 ubiquitylation after TLR stimulation.

**DISCUSSION**

Our previous studies showed that A20 restricts TLR-induced NF-κB signaling (22). Our current studies indicate that A20 regulates TRIF-dependent NF-κB signaling but not IRF3 signaling. Collectively, these data suggest that the major function of A20 in TLR signaling is to directly regulate a signaling event that is common to MyD88- and TRIF-dependent NF-κB signaling and that is not shared by TRIF-dependent IRF3 signaling. TRAF6 is an E3 ligase that is involved in both MyD88- and TRIF-dependent NF-κB signaling (28–30). Upon stimulation with TLR ligands, TRAF6 is recruited to the TLR in both MyD88-dependent and -independent signaling pathways. TRAF6 autoligates lysine-63 ubiquitin chains upon TLR or IL-1R stimulation to activate NF-κB, and our previous studies showed that A20 is able to use its OTU protease domain to deubiquitylate TRAF6 (22, 30). Thus, TRAF6 may be a physiological target for A20 during TLR signaling. To directly determine if A20 is essential for regulating TRAF6 ubiquitylation during TLR stimulation, A20+/+ and A20−/− BMDMs were stimulated with LPS for various periods of time, after which cell lysates were analyzed for their expression of ubiquitylated TRAF6. Endogenous TRAF6 is ubiquitylated after 20 min of LPS stimulation (Fig. 8). Importantly, whereas TRAF6 ubiquitylation is transiently induced in A20+/+ cells, declining after 40 min, TRAF6 ubiquitylation is prolonged to 40 and 60 min in A20−/− cells (Fig. 8). As these lysates were boiled in 1% SDS before immunoprecipitation of TRAF6, the ubiquitylated proteins we observe are likely to be predominantly ubiquitylated TRAF6 rather than noncovalently associated proteins. This finding indicates that A20 is essential for restricting endogenous TRAF6 ubiquitylation after TLR stimulation.
the intestinal epithelium. MyD88-dependent TLR signals may be beneficial for intestinal homeostasis under normal circumstances (2, 31). The appreciation of such homeostatic TLR signals has raised several questions, including whether homeostatic TLR signals are potentially inflammatory. Our finding that unperturbed A20−/− MyD88−/− mice are dramatically less inflamed and survive far longer than A20−/− MyD88+/+ mice suggests that homeostatic TLR signals can in fact be profoundly inflammatory and rapidly lethal. We have also elucidated a new function for A20 in regulating TRIF-dependent NF-κB signals but not IRF3 signals.

The idea that homeostatic MyD88 signals can cause spontaneous inflammation is broadly consistent with recent findings that constitutive MyD88 signals drive colitis and eventual mortality in IL-10−/− mice (3). However, although IL-10−/− mice develop progressive colitis during the first 4–6 mo of life and eventually die from this disease, the majority of A20−/− mice die within the first 3–6 wk of life from widespread systemic inflammation and cachexia (3, 32). Thus, our findings indicate that homeostatic MyD88 signals can drive severe perinatal systemic inflammation in addition to intestinal inflammation. Moreover, although IL-10 is particularly important for restricting intestinal inflammation, A20 is critical for restricting systemic inflammation.

Although A20 prevents widespread inflammation, a major source of constitutive, proinflammatory TLR signals in A20−/− mice may nevertheless be intestinal commensal bacteria. We have found that the removal of many of these bacteria with antibiotics reduced systemic inflammation in chimeric mice bearing A20−/− hematopoietic cells. Although previous studies have shown that intestinal bacteria can drive intestinal inflammation in several models (33), our current results link these bacteria and MyD88-dependent signals to A20-mediated regulation of innate immune cells and to systemic immune activation.

The different physiological roles of IL-10 and A20 in restricting MyD88-dependent signals is likely related to the distinct biochemical functions of these immunoregulatory proteins. IL-10 triggers STAT3-mediated transcriptional inhibition of proinflammatory genes, whereas our data indicate that A20 restricts MyD88-dependent TLR signals by regulating TRAF6 ubiquitylation. Thus, the critical physiological role that A20 plays in preventing constitutive homeostatic MyD88 signals from becoming inflammatory is likely caused by A20’s biochemical role in directly regulating MyD88-dependent signals.

Proteins such as A20, IRAK-M, ST2, and SIGIRR that restrict TLR signals could regulate the duration and/or intensity of TLR signals and modulate the cellular outcome of TLR signaling, thereby helping to determine whether TLR signals lead to homeostatic or inflammatory responses (14–18). Notably, although unperturbed mice lacking IRAK-M, ST2, and SIGIRR exhibit relatively modest inflammation and survive for longer than 9–12 mo of age, A20−/− mice typically die within the first few weeks of life (13–17). Although SOCS-1−/− mice exhibit perinatal lethality similar to A20−/− mice, SOCS-1−/− mice are rescued by the absence of IFN-γ signaling in SOCS-1−/− IFN-γ−/− double-mutant mice (19–21). Thus, among proteins known to directly restrict TLR signaling, A20 appears to play a more critical role in regulating homeostatic TLR-driven inflammation in vivo. There are several potential (non-mutually exclusive) reasons why A20 has a greater physiological impact on homeostatic TLR signals. First, A20 enzymatically regulates ubiquitylation of signaling proteins, thereby regulating both the activity and stability of such targets. Thus, A20 may not only deaminate signaling proteins but may also prevent further reactivation of these proteins. Second, A20 may regulate TLR signaling downstream of other known TLR regulating proteins, thus exerting a more definitive impact on the transcriptional outcome of TLR signals. Third, A20’s physiological roles in restricting MyD88-dependent TLR signals may be amplified by A20’s roles in regulating other non-TLR signals, e.g., TNF signals. However, marked survival differences between A20−/− MyD88−/− TNF−/− mice and A20−/− TNF−/− mice suggest that restricting MyD88 signals may be the dominant physiological function of A20 during homeostatic conditions. Finally, it is possible that A20 may also regulate IL-1 or IL-18 signals, which also use MyD88 (9). Although we have not yet

Figure 8. A20 is essential for restricting TRAF6 ubiquitylation. Immunoblotting analysis of endogenous TRAF6 ubiquitylation in LPS-stimulated BMDMs. A20−/− and A20+/+ BMDMs were stimulated with LPS, and cell lysates were generated at the indicated time points. Samples were boiled in 1% SDS to disassociate noncovalent protein–protein interactions. After dilution to 0.1% SDS, TRAF6 protein was immunoprecipitated from the lysates and immunoblotted for ubiquitin. Data are representative of at least three independent experiments.
evaluated the contribution of aberrant IL-1 or IL-18 signaling to spontaneous inflammation in A20\(^{-/-}\) mice, IL-1 receptor antagonist–deficient mice do not develop spontaneous inflammation within the first 4 mo of life (34). Thus, restricting basal IL-1 signals may not be critical for immune homeostasis in unperturbed mice. Collectively, A20 appears to be one of the most critical proteins for restricting homeostatic TLR signals in vivo.

Constitutive or basal MyD88–dependent signals could derive from microbial or host ligands. Our experiments with antibiotic-mediated depletion of commensal bacteria reinforce the notion that intestinal bacteria are a major source of MyD88–dependent signals triggering inflammation in unperturbed A20\(^{-/-}\) mice. Collectively with previous findings that constitutive MyD88–dependent signals from luminal bacteria are required for maintaining intestinal health, the current results suggest that a moderate but not excessive magnitude of MyD88 signaling in the intestine is essential for intestinal homeostasis.

In the intestine, homeostatic TLR signals might also be distinguished from inflammatory TLR signals by microenvironment- and cell type–specific TLR signals. For example, apical TLR signals on epithelial cells might preferentially induce homeostatic signals, whereas TLR signals on dendritic cells trigger inflammation. Although we have not yet directly addressed the roles of A20 in regulating epithelial cell function, our discoveries that radiation chimera bearing A20\(^{-/-}\) MyD88\(^{-/-}\) hematopoietic cells exhibit less spontaneous inflammation than chimeric mice bearing A20\(^{+/+}\) MyD88\(^{+/+}\) cells indicates that A20–mediated restriction of homeostatic TLR signals specifically in hematopoietic cells is important for immune homeostasis. Although it is possible that TLRs on epithelial cells may also be constitutively engaged on mucosal surfaces, our results using radiation chimera indicate that engagement of TLRs on hematopoietic cells occurs in unperturbed mice and must be properly restricted to maintain immune homeostasis. Indeed, one example of how TLRs on hematopoietic cells might tonically bind PAMPs from luminal microbes is via the extrusion of dendrites by mucosal dendritic cells through epithelial tight junctions into the intestinal lumen (35–37). Future studies using gene–targeted mice bearing lineage–specific deletions of A20 should facilitate studies of A20’s specific role in dendritic cells and other cell types.

The generation and characterization of A20\(^{-/-}\) MyD88\(^{-/-}\) mice also facilitates further studies of A20’s roles in regulating additional signaling pathways. Milder but progressive spontaneous inflammation in older double-mutant A20\(^{-/-}\) MyD88\(^{-/-}\) mice and triple-mutant A20\(^{-/-}\) MyD88\(^{-/-}\) TNF\(^{-/-}\) mice suggests that A20 regulates additional immune signals besides MyD88–dependent TLR and TNF signals. TRIF–dependent TLR signals may be one of these types of signals, and we have used A20\(^{-/-}\) MyD88\(^{-/-}\) mice to discover that A20 is also required for restricting TRIF–dependent TLR responses. We have also used A20\(^{-/-}\) MyD88\(^{-/-}\) and A20\(^{-/-}\) MyD88\(^{-/-}\) TNF\(^{-/-}\) cells to decipher the biochemical mechanism by which A20 regulates TLR signaling. Specifically, both A20\(^{-/-}\) MyD88\(^{-/-}\) mice and BMDMs derived from these mice produce more type I IFNs than A20\(^{+/+}\) MyD88\(^{-/-}\) mice and BMDMs after LPS stimulation. Previous work demonstrated that TRIF associates with TANK-binding kinase 1, TRAF3, and TRAF6 during the activation of NF-κB and IRF3 signaling (10, 38). Both NF-κB and IRF3 are required for TRIF–dependent IFN transcription. Receptor–interacting protein (RIP) 1 ubiquitylation, possibly mediated by TRAF6, is also involved in TRIF–dependent NF-κB signaling (39, 40). Our data suggest that A20 restricts TRIF–dependent NF-κB signaling and not IRF3 phosphorylation. Hence, A20’s nonredundant function in regulating TRIF–dependent signaling appears to be restricted to NF-κB signaling. A20 can inhibit TRIF signaling and can bind TRAF6, Nef–associated kinase/TANK-binding kinase 1, and IκB kinase ε (41–43). Thus, it is likely that A20 restricts TRIF–dependent signaling by directly regulating proteins in this signaling complex. Our data do not support the previous finding that heterologous A20 expression inhibits IRF3 dimerization, and these differences could be caused by the distinct experimental systems used (43). Collectively, our data suggest that A20 restricts TRIF–dependent IFN responses by limiting TRIF–dependent NF-κB signaling.

Previous work has shown that conjugation of K63–linked ubiquitin chains to TRAF6 is an essential step in TRM–induced NF-κB activation (30). Our earlier studies also showed that A20 is a ubiquitin–modifying enzyme that can both deconjugate ubiquitin chains and ligate ubiquitin onto target proteins such as RIP and TRAF6 (18, 23). Our current data directly demonstrate that A20 is essential for restricting endogenous TRAF6 ubiquitylation after TLR stimulation, further reinforcing this biochemical mechanism. As TRAF6 binds to both MyD88 and TRIF after TLR stimulation, TRAF6 likely activates NF-κB signals in both of these pathways. RIP1 is also involved in TRIF–dependent TLR signaling, and A20 restricts RIP1 ubiquitylation in response to TNF (39, 40). Thus, it is possible that A20 also restricts TRIF–dependent NF-κB signaling by restricting RIP1 ubiquitylation. However, RIP1 is not involved in MyD88–dependent NF-κB signaling, so it is unlikely that A20’s effects on RIP1 can explain all of A20’s roles in TLR signaling. In contrast, A20’s restriction of TRAF6 ubiquitylation may be a common mechanism by which A20 restricts both MyD88– and TRIF–dependent NF-κB signaling.

TRAF3 has recently been implicated in TRIF–dependent signaling (44, 45). TRAF3 appears to be more important for IRF signals, whereas TRAF6 is more important for NF-κB signaling. We and others have not been able to obtain evidence that A20 binds to or modifies TRAF3 (unpublished data), and our finding that IRF3 phosphorylation occurs normally in the absence of A20 suggests that A20 may preferentially regulate TRAF6– but not TRAF3–dependent signaling. These two proteins (along with TRAF2, TRAF4, and TRAF5) share structurally homologies, including RING and Zn finger motifs that may mediate E3 ligase activity. Selective regulation of TRAF6– but not TRAF3–dependent signaling by A20 might be explained if A20 preferentially binds TRAF6 rather than TRAF3. Alternatively, although TRAF6 is modified with K63–linked polyubiquitin chains during TLR–initiated signal transduction,
it is currently unclear if TRAF3 is similarly ubiquitylated. Thus, A20 may bind K63 polyubiquitylated TRAF6, whereas TRAF3 may not undergo this type of ubiquitylation modification. In either scenario, our findings indicate that A20 is a selective regulator of TRAF6- and not TRAF3-dependent signal transduction and provides new insights into how NF-κB and IRF signaling may be discriminated and differentially regulated.

Our finding that A20+/−/MyD88−/− mice virtually all survive to adulthood with modest amounts of inflammation contrasts sharply with our previous findings that A20−/−, A20−/−/RAG−1−/−, A20−/−/TNF−/−, and A20−/−/TNFR1−/− mice all spontaneously develop severe inflammation, cachexia, and premature death (18, 19). This result suggests that dysregulated homeostatic TLR signals stimulate downstream innate and adaptive immune signals in the absence of A20 (e.g., TNF, IL-12, IL-6, and chemokine production, T cell activation, B cell activation, etc.). Although A20 may play multiple important roles in regulating immune cell signals, its role in restricting homeostatic TLR signals may be physiologically critical because TLR signals are situated at the apex of immune responses. Moreover, our results showing that antibiotics ameliorate inflammation in A20−/−/MyD88−/− chimeric mice suggest that commercial intestinal flora trigger these homeostatic MyD88−/− dependent TLR signals. Finally, we have established a biochemical mechanism by which A20 restricts both MyD88−/− and TRIF−dependent TLR signals. In summary, our findings demonstrate the profoundly pro-inflammatory nature of homeostatic MyD88−/− dependent signals and identify A20 as a critical protein that prevents homeostatic signals from becoming inflammatory. Further studies elucidating how A20 restricts different types of TLR signals should yield important insights into the complex interplay between commensal microbes and host immune cells, as well as the biochemical mechanisms by which signals triggered by microbial molecules are regulated and interpreted.

### Materials and Methods

#### Mouse and cell preparations.

The generation and characterization of A20−/− mice have been previously described (20). A20−/−/Ly5.2 mice were backcrossed for eight generations onto a C57BL/6J background before being interbred with R. Medzhitov (Yale University, New Haven, CT) and were backcrossed for five generations onto a C57BL/6J background. MyD88−/− mice were obtained from K. Harada (University of Virginia, Charlottesville) and were backcrossed for eight generations onto a C57BL/6J background. MyD88−/− mice were generated by transfer of wild-type or A20−/− bone marrow HSCs into lethally (700 rads) congenic Ly5.2+ C57BL/6j recipient mice. All mice were given trinitrophenyl-sulfamethoxazole immediately after reconstitution, 2 wk after reconstitution, some mice were given 0.5 grams per liter of vancomycin hydrochloride (Novaplan), 1 gram per liter of ampicillin (Sandoz), 1 gram per liter of neomycin sulfate (Pharma-Tek), and 1 gram per liter of metronidazole (Baxter) in drinking water. After 3–4 wk, mice were killed and analyzed by flow cytometry. To quantitate colonic microflora, fecal matter was removed from colons, weighed, and homogenized in luria broth at 0.1 g/ml before culturing on LBagar plates.

#### TLR ligands and ELISAs.

LPS from *S. typhimurium* and poly (I:C) (GE Healthcare) were used at concentrations of 1 μg/ml and 50 μg/ml, respectively. BMDMs were stimulated with either LPS or poly (I:C) for 24 h, and culture supernatants were assayed by ELISA for IFN-γ, TNF-α, IL-6, and chemokine production, T cell activation, B cell activation, etc.). Although A20 may play multiple important roles in regulating immune cell signals, its role in restricting homeostatic TLR signals may be physiologically critical because TLR signals are situated at the apex of immune responses. Moreover, our results showing that antibiotics ameliorate inflammation in A20−/−/MyD88−/− chimeric mice suggest that commercial intestinal flora trigger these homeostatic MyD88−/− dependent TLR signals. Finally, we have established a biochemical mechanism by which A20 restricts both MyD88−/− and TRIF−dependent TLR signals. In summary, our findings demonstrate the profoundly pro-inflammatory nature of homeostatic MyD88−/− dependent signals and identify A20 as a critical protein that prevents homeostatic signals from becoming inflammatory. Further studies elucidating how A20 restricts different types of TLR signals should yield important insights into the complex interplay between commensal microbes and host immune cells, as well as the biochemical mechanisms by which signals triggered by microbial molecules are regulated and interpreted.

#### RNA isolation and quantitative PCR analyses.

RNA was isolated from spleens using extraction of homogenized tissue (RNeasy Mini Kit; QIAGEN). RNA was treated with RNase-free DNase (QIAGEN) to eliminate contaminating genomic DNA and quantified using a spectrophotometer (ND-1000, Nanodrop). Complementary DNA was generated using an RT kit (Quantitect; QIAGEN), and real-time quantitative PCR was performed using an SYBR green PCR kit (Quantitect; QIAGEN). The following gene-specific primers were used for quantitating TNF-α (sense, 5′-TGGCCCTCCCTCTCATCACAGTT-3′; antisense, 5′-TCTTCACCACCTTGGTGTTCG-3′), IL-1β (sense, 5′-CCGGTACCTTCTCCAGATTGA-3′; antisense, 5′-GGGACACTGACACACACACACA-3′), IL-6 (sense, 5′-AAAACCTTCTTATGGAGATGCAGTG-3′; antisense, 5′-TTGAGGGCCTTTGATGAGTA-3′), β-actin (sense, 5′-AAGTTGACGTGAGCATTTCTTA-3′; antisense, 5′-TGCCTGTTATCTAATGCTT-3′), and GAPDH (sense, 5′-AACGGGAGAAGACTGTTGTC-3′; antisense, 5′-TGTTTGAATTTCACTCTCCCTCCACTGCG-3′)

#### Immunoblotting and immunoprecipitation.

In vitro TLR signal analyses were performed with antibodies specific for F4/80 (Serotec) and Mac-1 (BD Biosciences). Macrophage purity was typically 90–95%. BMDMs were allowed to adhere to tissue culture plastic overnight before stimulation.
REFERENCES


