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Polyubiquitin binding to ABIN1 is required to prevent autoimmunity

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The protein ABIN1 possesses a polyubiquitin–binding domain homologous to that present in nuclear factor κB (NF-κB) essential modulator (NEMO), a component of the inhibitor of NF-κB (IκB) kinase (IKK) complex. To address the physiological significance of polyubiquitin binding, we generated knockin mice expressing the ABIN1[D485N] mutant instead of the wild-type (WT) protein. These mice developed all the hallmarks of autoimmunity, including spontaneous formation of germinal centers, isotype switching, and production of autoreactive antibodies. Autoimmunity was suppressed by crossing to MyD88−/− mice, demonstrating that toll–like receptor (TLR)–MyD88 signaling pathways are needed for the phenotype to develop. The B cells and myeloid cells of the ABIN1[D485N] mice showed enhanced activation of the protein kinases TAK, IKK-α/β, c-Jun N-terminal kinases, and p38 mitogen-activated protein kinase and produced more IL-6 and IL-12 than WT. The mutant B cells also proliferated more rapidly in response to TLR ligands. Our results indicate that the interaction of ABIN1 with polyubiquitin is required to limit the activation of TLR–MyD88 pathways and prevent autoimmunity.
may induce a conformational change that facilitates phosphorylation of the activation loop of the canonical IKKs by TAK1 and/or autophosphorylation.

The canonical IKKs activate NF-κB by phosphorylating the inhibitory IκBα component of this transcription factor, which marks IκBα for K48-linked polyubiquitylation by the SCFTRCP E3 ligase and proteasomal destruction. The canonical IKKs also switch on the protein kinase Tpl2 by phosphorylating its inhibitory p105/NF-κB1 component. Similarly, TAK1 not only initiates activation of thecanonical IKKs but is also required to activate c-Jun N-terminal kinase (JNK) and p38α mitogen-activated protein kinase (MAPK) via the TLR–MyD88 signaling pathway. The canonical IKKs and MAPKs then catalyze many further phosphorylation events that control the transcription, translation, processing, and secretion of inflammatory mediators (Sato et al., 2005; Shin et al., 2005).

Interestingly, the polyubiquitin-binding domain in NEMO, originally termed A20-binding inhibitor of NF-κB (ABIN) homology domain 2 (AHHD2; Heyninck et al., 2003), but later renamed the ubiquitin-binding domain in ABIN and NEMO (UBAN; Wagner et al., 2008), is found in four other human proteins, termed NRP (NEMO-related protein, also called optineurin), ABIN1, ABIN2, and ABIN3. The ABINs are so named because they were originally identified in a yeast two-hybrid screening using the protein deubiquitylase A20 as bait and because they were found to inhibit NF-κB–dependent gene transcription when overexpressed in cells (Heyninck et al., 1999).

Recently, ABIN1 knockout mice were generated and characterized (Oshima et al., 2009). These mice were found at normal Mendelian ratios up to embryonic day (E) 18.5, but the embryos were smaller and more anemic than WT embryos and died during late embryogenesis from fetal liver apoptosis and hypoplasia. Embryonic fibroblasts isolated from the ABIN1−/− mice were hypersensitive to TNF-induced programmed cell death and the lethality could be rescued by crossing to mice that did not express the TNFR1 receptor (Oshima et al., 2009).

To understand the physiological role of the polyubiquitin-binding function of ABIN1, we generated knockin mice that express the ABIN1[D485N] mutant instead of the WT protein, a mutation which is equivalent to the polyubiquitin-binding-defective NEMO[D311N] mutant described already. Unexpectedly, these knockin mice had a quite different phenotype from the ABIN1−/− mice. They were born at normal Mendelian frequencies and the adults were of normal size and weight. However, they then developed a lupus-like autoimmune disease, which could be prevented by crossing to MyD88-deficient mice. These and other results presented in this paper demonstrate that the interaction of ABIN1 with polyubiquitin chains limits the strength of signaling downstream of TLR–MyD88 and that this is critical to prevent autoimmunity.

RESULTS

Interaction of ABIN1 with polyubiquitin chains in vitro and generation of ABIN1[D485N] knockin mice

The mutation of Asp311 of NEMO to Asn prevents it not only from binding to K63-pUb chains (Ea et al., 2006; Wu et al., 2006) but also from binding to linear polyubiquitin chains (Rahighi et al., 2009). Linear polyubiquitin chains are thought to be generated by the LUBAC E3-ligase (Kirisako et al., 2006) and to
ABIN1[D485N] mutant was unable to bind to polyubiquitin chains in vivo, we immunoprecipitated the endogenous ABIN1 from BMDM extracts, followed by immunoblotting with anti-IRAK1 (to detect polyubiquitylated-IRAK1) and anti-ubiquitin. These experiments showed that stimulation with LPS induced the binding of ABIN1 to polyubiquitylated IRAK1 (Windheim et al., 2008) but not to unmodified IRAK1. In contrast, the ABIN1[D485N] mutant did not capture polyubiquitylated IRAK1 (Fig. 1 C).

Gross analysis of ABIN1[D485N] knockin mice

The ABIN1[D485N] mice were the same size and weight as WT littermates (Fig. 2 A) and were born at the expected Mendelian frequencies (not depicted). However, examination of their internal organs at 2–3 mo of age revealed enlarged spleens which, by 4 mo, were four to five times the weight of those from WT mice (Fig. 2 A). Both the white pulp and the red pulp were greatly enlarged in the spleen, although the splenic architecture was maintained (Fig. 2 B).

The ABIN1[D485N] mutant was expressed at similar levels to WT ABIN1 in BM-derived macrophages (BMDMs), B cells, and T cells (Fig. S1, C–E). To check that the mouse ABIN1[D485N] mutant was unable to bind to polyubiquitin chains in vivo, we immunoprecipitated the endogenous ABIN1 from BMDM extracts, followed by immunoblotting with anti-IRAK1 (to detect polyubiquitylated-IRAK1) and anti-ubiquitin. These experiments showed that stimulation with LPS induced the binding of ABIN1 to polyubiquitylated IRAK1 (Windheim et al., 2008) but not to unmodified IRAK1. In contrast, the ABIN1[D485N] mutant did not capture polyubiquitylated IRAK1 (Fig. 1 C).

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ABIN1[D485N] knockin mice had significantly enlarged LNs (Fig. 2 C) and 60–70% of the knockin mice had large nodule-like structures on the intestinal wall, which were identified as Peyer’s Patches (Fig. 2 D and E). Although young ABIN1[D485N] mice appeared to be superficially healthy, by 20 wk they all developed signs of severe autoimmunity, resulting in the majority of the knockin mice being terminated by 6 mo of age. This phenotype was exacerbated in the female mice during pregnancy.
the proportions of different B cell populations in the spleen, LNs, and peritoneal cavity of the ABIN1[D485N] mice did not differ significantly from those in the WT mice (Figs. S2, B–E). We also found that the ABIN1[D485N] knockin mice had four to five times the normal level of monocytes in the blood (unpublished data).

T cell development in the thymus was normal (Fig. S2 F). In contrast, analysis of the splenic and LN T cell populations demonstrated an increased proportion of activated effector T cells (CD62Llo and CD44hi) and a reduced proportion of naive T cells (CD62Lhi) compared with age-matched controls (Fig. 3 D).

Defects in immune cells with spontaneous germinal center formation in the ABIN1[D485N] knockin mice

To investigate immunological changes in the mice, we performed flow cytometry analyses of various lymphoid organs. Analysis of splenic and LN cells revealed more B cells and granulocytes (CD11b+ Gr-1+) in the ABIN1[D485N] mice compared with WT mice (Fig. 3, A and B; and Fig. S2 A). The expression of MHCII and CD86 receptors on B cells was also increased in the spleen (Fig. 3 C) and LNs (not depicted) of the ABIN1[D485N] mice, indicating that the B cells were activated. However, the development of B cells in the BM and

Figure 3. Defects in immune cells with spontaneous germinal center formation in ABIN1[D485N] mice. (A–F and H) Flow cytometric profiles of spleen (SPL) and LNs. Profiles were gated on: lymphocytes by FSC/SSC (C–F and H); live cells (A); TCR-β+ CD4+ (D and E), and B220+ (C and F). The numbers within figures indicate the percentages of different cells. GC B, germinal center B cell. (A) Flow cytometric dot plots of spleen (top) and LN (bottom) showing frequencies of the CD11b+ Gr-1+ (macrophages and granulocytes) population in ABIN1[D485N] and WT mice. (B) Total cell numbers in spleens of ABIN1[D485N] mice compared with WT. Error bars show the mean ± SEM. ***, P < 0.001 (two-tailed Student t test). (C) Expression of B cell activation markers from WT (shaded area) and ABIN1[D485N] (black line). (D–F) Contour plots (left) and graphical analysis (right) showing percentages of cell populations in spleen (top) and LN (bottom). Each symbol represents one mouse and the horizontal bars show the mean of the values obtained. (D) Activated (CD4+ CD44hi CD62Llo) and naive (CD4+ CD62Lhi) T cells. (E) Tfh cells (CD4+ CXCR5+ PD-1hi). (F) Germinal center B cells (B220+ GL-7+ CD95+). (G) Immunohistochemistry of spleens from 16-wk-old mice with germinal centers (brown) stained with peanut agglutinin (PNA). Bars, 0.5 mm. (H) Expression of CD138 and B220 in spleen showing plasma cells in ABIN1[D485N] mice and WT mice. All mice analyzed were 12–16 wk old, and data are representative of at least three independent experiments with three to four mice of each genotype (A–F and H) or single experiment with six mice per genotype (G).
Elevated levels of immunoglobulins and autoantibodies in the serum of ABIN1[D485N] mice

The hyperplasia of Tfh cells and germinal center B cells in the spleen and LN led us to investigate the serum immunoglobulins levels. A variety of immunoglobulin isotypes were elevated in the ABIN1[D485N] knockin mice compared with age-matched WT mice, which included both T cell–dependent and T cell–independent immunoglobulins. In particular, the levels of pathogenic immunoglobulins were increased significantly after 20 wk. The circulating levels of antinuclear antibodies and antibodies against double-stranded (ds) DNA were also increased significantly after 16 wk or 20 wk in the ABIN1[D485N] mice, demonstrating that some of the antibodies in the knockin mice were being pathogenic.

However, the proportion of CD4+ and CD8+ T cells in the spleen was normal (Fig. S2 G).

It has been reported that an increased number of follicular helper T cell (Tfh) cell is associated with autoimmunity, and we therefore studied whether Tfh cell number was altered in the ABIN1[D485N] mice. These experiments revealed that the proportion of Tfh cells (CD4+ PD1hi and CXCR5+) in the spleen and LN was greatly increased, as were the number of germinal center B cells (B220+ CD95+ and GL7+; Fig. 3 F). Consistent with more germinal center B cells, there was increased formation of germinal centers in the spleens of ABIN1[D485N] mice (Fig. 3 G). There was also an increase in the proportion of extrafollicular plasma cells in the spleen of ABIN1[D485N] mice (Fig. 3 H).

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produced against self-cellular components, a feature of autoimmune disease. The heterozygous ABIN1\[D485N\]+/- mice, which did not develop any pathological abnormalities, only had marginally elevated levels of anti-dsDNA (Fig. 4 B).

High levels of pathogenic immunoglobulins in the serum are known to be deposited in the kidney and the blood vessels, initiating an inflammatory reaction in these organs. We found that there was immune complex deposition in the kidney of the ABIN1\[D485N\] mice, leading to activation of the complement pathway (Fig. 4 D) and the development of severe renal disease at 20–24 wk of age. Histologically, this was revealed by severe generalized global membranoproliferative glomerulonephritis of the kidney with infiltration of neutrophils and plasma cells (Fig. 4 E, top). There was also thickening of glomerular capillary loops and Bowman’s capsule basement membrane regions by periodic acid-Schiff (PAS)–positive material (Fig. 4 E, bottom).

Vascular lesions were noted at the same age in multiple sites, including splenic arterioles (Fig. 4 F) and heart-base arteries (Fig. 4 G), with occasional involvement of Peyer’s Patch arterioles and peripancreatic arteries (not depicted). The normal architecture of the walls of affected splenic arterioles was largely replaced by fibrinous material with scattered nuclear debris and mononuclear inflammatory cells (fibrinoid necrosis), and by extravascular leakage (Fig. 4 F). Similar changes were noted in heart-base arteries but with more severe inflammatory cell infiltrates, including neutrophils, macrophages, and reactive fibroblasts, which not only segmentally obliterated arterial wall architecture but also extended into the adjacent atrial myocardium (Fig. 4 G).

The combination of lesions may explain why all the ABIN1\[D485N\] mice developed severe symptoms of autoimmune disease, requiring them to be culled within 6 mo, if they had not died already. In contrast, no WT control mice had died at this age.

**Enhanced proliferation, cytokine production, and activation of signaling pathways in B cells and myeloid cells**

Enhanced activation of B cells and myeloid cells in ABIN1\[D485N\] mice. (A) Naive purified splenic B cells from WT and ABIN1\[D485N\] mice were stimulated with 200 ng/ml LPS, 10 µg/ml LTA, 200 ng/ml of the TLR7 agonist R848, 10 µg/ml α-IgM, or 1 µg/ml α-CD40 for 72 h or left unstimulated (control) before pulsing for a further 16 h with [3H]thymidine (0.5 µCi/well). [3H]thymidine incorporation into DNA was measured by harvesting and washing the cells followed by measurement of radioactivity incorporated. (B) Flow cytometric analysis of surface expression of the activation marker CD86 after stimulation of purified B cells with the agonists indicated. The filled histograms show results for WT cells and the empty histograms show the ABIN1\[D485N\] cells. (C) IL-6 and IL-12p40 secreted into the culture medium of B cells 48 h after exposure to the agonists indicated. (D) IL-6 and TNF secreted into the culture medium of BMDC 24 h after exposure to 100 ng/ml LPS, 2 µg/ml LTA, 1 µg/ml Pam3CSK4, 1 µg/ml R848, or 1 µM of the TLR9 agonist ODN 1826. Data are representative of five (A) or three (B–D) independent experiments with three to four mice of each genotype analyzed together. Error bars represent mean ± SD. *, P ≤ 0.05; **, P ≥ 0.005; ***, P ≤ 0.005 (two-tailed Student’s t test).
more marked in the ABIN1[D485N] knockin cells (Fig. 5 B), and this might contribute to the increased activation of T cells via interaction with the CD28 receptor on T cells.

In contrast, stimulation of purified CD4+ T cells with anti-CD3, alone or in combination with either CD28 or IL-2, resulted in similar proliferation rates in WT and ABIN1[D485N] knockin cells (Fig. S3 A), and the production of T cell cytokines tested was similar in different genotypes (Fig. S3 B). There was no effect of the knockin mutation on the expression of inducible co-stimulator (ICOS) or the T cell activation markers CD25 or CD69 after TCR stimulation (unpublished data), again indicating that the increased proportion of activated T cells and Tfh cells may not be explained by a T cell–intrinsic defect. Collectively, these results indicated that the T cell phenotypes observed in the ABIN1[D485N] mice may not have been caused by an intrinsic alteration in T cell receptor signaling. It has been well documented that regulatory T cells (T reg cells) play an important role in autoimmunity by limiting the activation of T cells, and that a decreased T reg cell number can cause autoimmunity. However, the proportion of T reg cells in the

the mutant mice because these cells are major producers of immunoglobulins in vivo. Using purified naïve B cells from the spleens of 5–6-wk-old mice before the onset of splenomegaly, we found that the TLR4 agonist LPS, the TLR2/6 agonist lipoteichoic acid (LTA), and the TLR7 agonist R848, as well as anti-IgM and anti-CD40, stimulated B cell proliferation more strongly in the ABIN1[D485N] mice than in the WT mice (Fig. 5 A). In contrast, the B cells from WT and ABIN1[D485N] mice proliferated similarly in response to the TLR9 ligand ODN1826 (unpublished data). The increased proliferation of B cells in response to anti-IgM and anti-CD40 was not explained by differences in the expression of these receptors, which were similar in ABIN1[D485N] and WT mice (unpublished data). The stimulation of B cells with either TLR agonists or anti-CD40 also increased the expression of the co-stimulatory molecule CD86, which was
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ABIN1[D485N] mice was only slightly higher than that in WT mice (Fig. S3 C).

The stimulation of B cells with LPS alone, or the combination of LPS with anti-CD40, also induced higher levels of IL-6 and IL-12p40 secretion in the ABIN1[D485N] mice compared with WT mice. Similar results were found with LTA and R848 (Fig. 5 C). Moreover, TLR stimulation of other immune cells of myeloid origin, such as BM-derived DCs (BMDCs), also increased production of IL-6 and other proinflammatory cytokines, such as TNF, more markedly in the cells from the ABIN1[D485N] mice (Fig. 5 D).

The results described in the previous paragraph suggested that signaling events required for the production of cytokines by B cells and myeloid cells were being switched on more strongly in the ABIN1[D485N] mice compared with WT, and they led us to examine the activation of these pathways. Stimulation of naive splenic B cells with the TLR7 ligand R848 (Fig. 6 A), the TLR4 ligand LPS (Fig. S4 A), the B cell receptor (BCR) ligand (anti-IgM; Fig. 6 B), or the CD40 agonist (anti-CD40; Fig. 6 C) caused enhanced phosphorylation (activation) of the canonical IKKs (IKK-α/β) and the phosphorylation of their substrates p105/NF-κB1 and IκBα in the ABIN1[D485N] mice compared with WT mice, and, consistent with enhanced phosphorylation of IκBα, there was a more rapid proteasomal degradation of this protein. There was also modestly enhanced phosphorylation (activation) of the MAPKs JNK1/2 and p38MAPK. Enhanced activation of the canonical IKKs and MAPKs was also observed in BMDC and BMDM from ABIN1[D485N] mice compared with WT mice after stimulation with LTA or the TLR7 agonist R848 (Fig. 6, D and E; and Fig. S4, B and C).

The finding that activation of MAPKs, as well as the canonical IKK complex, was enhanced in immune cells from the ABIN1[D485N] mice after TLR stimulation indicated that ABIN1 must be exerting its effect by suppressing the activation of a more upstream component of the pathway. This led us examine the activation of TAK1 and to show that it is also enhanced in BMDC from the knockin mice compared with WT mice (Fig. 6 F).

Enhanced activation of the nucleotide oligomerization domain (NOD) 1–NOD2 signaling pathway

The cytoplasmic proteins NOD1 and NOD2 detect peptidoglycan components from intracellular bacteria that have evaded the TLR defense system (Inohara et al., 2005). The activation of these receptors induces their interaction with the protein kinase RIP2, leading to the activation of NF-κB and IκBα in the ABIN1[D485N] mice compared with WT mice, and, consistent with enhanced phosphorylation of IκBα, there was a more rapid proteasomal degradation of this protein. There was also modestly enhanced phosphorylation (activation) of the MAPKs JNK1/2 and p38α MAPK. Enhanced activation of the canonical IKKs and MAPKs was also observed in BMDC and BMDM from ABIN1[D485N] mice compared with WT mice after stimulation with LTA or the TLR7 agonist R848 (Fig. 6, D and E; and Fig. S4, B and C).
(not depicted) induced a far stronger activation of IKK-α/β and MAPKs in BMDM from the ABIN1[D485N] mice than WT mice.

The absence of MyD88, but not RIP2, suppresses the autoimmune phenotype of ABIN1[D485N] mice

The results presented in the previous section suggested that enhanced activation of one or more cell signaling pathways might underlie or contribute to the autoimmune phenotype of the ABIN1[D485N] mice. To investigate the importance of the TLR–MyD88–dependent and NOD1/2–RIP2 signaling pathways, we crossed the ABIN1[D485N] mice with mice that do not express the adaptor MyD88 (required for TLR–MyD88 signaling) signaling or with mice that do not express RIP2. The ABIN1D485N)xRIP2−/− mice still displayed all the gross phenotypes of the ABIN1[D485N] mice (splenomegaly, lymphadenopathy, and intestinal nodules) and flow cytometry analysis did not show rescue of any immune phenotype (unpublished data). The ABIN1[D485N]xMyD88−/− mice, however, did not develop splenomegaly (Fig. 7 A), enlarged LNs, and intestinal nodules (not depicted). They had greatly reduced numbers of germinal center B cells in the spleen (Fig. 7 B), and serum levels of anti-dsDNA, antinuclear antibodies (Fig. 7 C), and different immunoglobulins (not depicted) induced a far stronger activation of IKK-α/β and MAPKs in BMDM from the ABIN1[D485N] mice than WT mice.

CD40 is not thought to signal via MyD88 (He et al., 2010a), but using B cells from age-matched ABIN1[D485N] mice, MyD88−/− mice, ABIN1[D485N]/MyD88−/− mice, and WT mice, we found that the enhanced activation of the canonical IKKs and JNK observed in CD40-stimulated B cells from the ABIN1[D485N] mice was reduced in B cells from the ABIN1[D485N]/MyD88−/− mice to the same level found in either WT mice or MyD88−/− mice (unpublished data). This suggests that the over-reactivity of the ABIN1[D485N]-expressing B cells to CD40 depends on MyD88 signaling. This suggests that there may be cross talk between the CD40 and MyD88 signaling pathways, which would be interesting to investigate in future studies.

Generation of ABIN1[D478–606] mice

The way in which the targeting vector for the ABIN1[D485N] mice was designed allowed the targeted region to be removed by crossing to CRE deleter mice, which generated a form of ABIN1 in which residues 478–606 were deleted. This deletion did not cause the truncated protein to misfold and be degraded, and the ABIN1[D478–606] mutant was actually expressed at higher levels than WT ABIN1 in BMDM (Fig. S6 A) and embryonic fibroblasts (not depicted). The ABIN1[D478–606] mice were also born at near Mendelian frequencies and had a similar phenotype to the ABIN1[D485N] mice (Fig. S6 B), although it was slightly more severe, in as much as the mice became moribund about a month earlier than the ABIN1[D485N] mice. As found in the ABIN1[D485N] mice, there was enhanced activation of TLR signaling pathways in BMDM from the ABIN1[D478–606] mice (Fig. S6 A).

TNF-induced apoptosis in embryonic fibroblasts from ABIN1[D485N] mice

ABIN1−/− mice were reported to be found at normal Mendelian ratios up to E18.5, but the embryos then died during late embryogenesis from fetal liver apoptosis and hypoplasia. Embryonic fibroblasts isolated from the ABIN1−/− mice were hypersensitive to TNF-induced apoptosis, and the lethality could be rescued by crossing to mice that did not express the TNFR1 receptor (Oshima et al., 2009). The TNF-stimulated apoptosis could be restored to normal levels by reintroducing WT ABIN1 but not by reintroduction of the ABIN1[D485N] mutant. Consistent with these findings, we found that embryonic fibroblasts from the ABIN1[D485N] knockin mice also showed enhanced TNF-induced apoptosis (Fig. S7, C and D). The reason why the ABIN1[D485N] and the ABIN1[D478–606] mice that we generated did not display embryonic lethality is unclear but could be the result of a difference in the background of the mice.

In contrast to the effect of TNF on apoptosis, the TNF-stimulated activation of NF-κB was reported to be similar in embryonic fibroblasts from ABIN1−/− mice and WT mice (Oshima et al., 2009). Consistent with this finding, we did not see enhanced activation of MAPKs or the canonical IKK complex in embryonic fibroblasts from the ABIN1[D485N] mice, although these signaling pathways were activated a little more strongly in IL-1–stimulated fibroblasts from the ABIN1[D485N] mice (Fig. S7).

DISCUSSION

In this paper, we demonstrate that replacing a single aspartate in the UBAN of ABIN1 (Asp485) by an asparaginyl residue, which suppresses the binding of ABIN1 to K63-pUb chains or linear pUb chains, causes autoimmunity in mice. Immune cells from the ABIN1[D485N] knockin mice showed enhanced activation of signaling pathways in response to TLR agonists that signal via MyD88, leading to increased B cell proliferation and enhanced production of IL-6 and IL-12p40 production by B cells and elevated IL-6 and TNF by myeloid cells. The critical role of TLR signaling pathways in driving autoimmunity was indicated by the finding that the phenotype was suppressed completely when the ABIN1[D485N] knockin mice were crossed to MyD88−/− mice. These observations raised the question of how increased signaling via TLR–MyD88 pathways can lead to autoimmunity.

Autoimmunity probably results from the formation of many germinal centers that were present in the spleens of the ABIN1[D485N] mice and are responsible for antibody isotype switching and the production of pathogenic antibodies. As suggested by others (Vinuesa et al., 2005, 2009), the formation
of germinal centers may be triggered by the presence of abnormally high numbers of Tfh cells, which was a feature of the spleen and LNs of the ABIN1[D485N] knockin mice. An increase in the number of Tfh cells can result from the aberrant expression of ICOS molecules on T cells (Vinuesa et al., 2005) and be stimulated by IL-6 and IL-12 (Nurieva et al., 2008; Ma et al., 2009; Schmitt et al., 2009). We did not observe any changes in T cell receptor activation or increased production of ICOS by T cells in the ABIN1[D485N] mice compared with WT mice after TCR stimulation. It therefore seems more likely that increased IL-6 and IL-12 secretion in B cells and myeloid cells via the TLR–MyD88 pathway, coupled with increased expression of co-stimulatory molecules (CD80 and CD86) by antigen-presenting cells, underlies the increased number of Tfh cells in the mutant animals. Enhanced TLR signaling could, however, lead to germinal center formation in other ways. For example, the migration of B cells to form new germinal centers can be driven by stimulation of TLR4 (Hwang et al., 2009) or TLR7 (Bessa et al., 2010) in vivo, even in IL–21−/− mice where Tfh cell function is defective. Another potentially important mechanism by which TLR–MyD88 signaling pathways contribute to autoimmunity is via the hyperactivation of T reg cells, leading to loss of their ability to suppress effector T cell responses (Pasare and Medzhitov, 2004; Peng et al., 2005; Sutmuller et al., 2007). In summary, our studies with the ABIN1[D485N] mice provide striking support for the concept that abnormally high TLR–MyD88 signaling can trigger autoimmunity (Marshak-Rothstein, 2006).

The increased production of interferon α in plasmacytoid DCs by TLR7 and/or TLR9 agonists is reported to be involved in the development of lupus-like autoimmunity (Banchereau and Pascual, 2006), raising the question of whether ABIN1 prevents autoimmunity by limiting TLR7/TLR9-stimulated interferon α production by pDC. However, we were unable to detect any increase in interferon α production by pDCs from ABIN1[D485N] mice compared with pDCs from WT mice in response to the TLR7 agonist poly(dU) or the TLR9 agonist ODN1826 (unpublished data).

An important unresolved question is the identity of the endogenous ligands that initiate activation of the TLR–MyD88 pathways in these animals in vivo. One possibility is that endogenous host RNA and/or DNA is able to activate TLR7 and TLR9 in the ABIN1[D485N] mice but not in WT mice. Alternatively, components of commensal bacteria may activate the TLR–MyD88 pathways resulting in autoimmunity, as observed in mice that do not express TANK (TRAF-associated NF-κB activator), which also display enhanced activation of NF-κB in response to ligands that signal via the TLR–MyD88 dependent pathway (Kawagoe et al., 2009). To investigate whether commensal bacteria were involved in the development of the phenotype, we fed the mice from birth with broad spectrum antibiotics, but this had no effect on the phenotype. The level of anti-dsDNA in the serum after 16 wk was 20.47 ± 3.69 kU/ml (n = 8) and 19.42 ± 8.19 kU/ml (n = 9), respectively, in control and antibiotic-fed WT mice, and rose to 61.54 ± 33.26 kU/ml (n = 8) and 78.55 ± 31.57 kU/ml (n = 5), respectively, in control and antibiotic-fed ABIN1[D485N] mice. Moreover, the antibiotic-fed ABIN1[D485N] mice had similar enlargement of the spleen and LNs. In contrast, the control IL–10−/− mice, which were fed the same combination of antibiotics in parallel, were protected against the development of ulcerative colitis and anal prolapse, as expected.

How polyubiquitin binding to WT ABIN1 limits the strength of TLR–MyD88 signaling is a complex issue and a full molecular understanding will require further analysis. However, the studies we have performed in this paper indicate that ABIN1 exerts its inhibitory effects downstream of MyD88 but upstream of TAK1, explaining why the activation of JNK and p38α MAPK, as well as the canonical IKKs, is enhanced in immune cells from the ABIN1[D485N] mice. Enhanced activation of TAK1 can also explain why the NOD1/2–RIP2 signaling pathway was activated more strongly in peptidoglycan-stimulated BMDM from ABIN1[D485N] mice and why B cells from the mutant mice proliferated more rapidly than B cells from the WT mice in response to BCR and CD40 agonists because TAK1 is known to be required for NOD1/2–RIP2 (Windheim et al., 2007), BCR, and CD40 signaling (Sato et al., 2005). ABIN1 was originally identified as a protein that interacts with the deubiquitylase A20. Like ABIN1, A20 is known to function as a negative regulator of the innate immune system and mice with a conditional knockout of A20 in B cells develop autoimmunity (Boone et al., 2004; Tavares et al., 2010; Chu et al., 2011). Moreover, human polymorphisms in A20 predispose to autoimmune diseases (Plenge et al., 2007; Thomson et al., 2007; Nair et al., 2009). A priori, one might therefore have imagined that by binding to K63-pUb chains/proteins generated in response to TLR activation (see Introduction), ABIN1 may recruit A20 to its substrates, facilitating the A20-catalysed hydrolysis of K63-pUb chains/proteins and the inhibition of TLR–MyD88–dependent signaling. Cells expressing the polyubiquitin binding–defective ABIN1[D485N] mutant would be unable to perform this function, leading to increased levels of K63-pUb chains/proteins and enhanced recruitment and activation of the TAK1 complex. However, whether this is the way in which ABIN1 limits the strength of signaling is unclear for several reasons. First, the bacterially expressed A20 catalytic domain cleaves K48-pUb chains and not K63-pUb chains or linear polyubiquitin chains in vitro (Komander et al., 2009), although A20 may be able to hydrolyze K63-pUb chains when overexpressed in mammalian cells (Wertz et al., 2004), in which case there may be a mechanism for altering the specificity of A20 in vivo. Second, a truncated form of ABIN1, ABIN1[444–601], which does not interact with A20, inhibits TNF-stimulated NF-κB–dependent gene transcription as effectively as WT ABIN1 in overexpression experiments (Heyninck et al., 2003). Third, the level of A20 is low in unstimulated cells and increases greatly after stimulation for 1 h with TLR agonists (Fig. 6, C and D), but the enhanced activation of signaling pathways in immune cells from the ABIN1[D485N] mice can be observed...
after only 10–15 min (Fig. 6), demonstrating that this effect is independent of the induction of A20. Fourth, although the B cell–specific knockout of A20 leads to autoimmunity, the onset of the pathology seems to be delayed by several months compared with the ABIN1[D485N] mice (Tavares et al., 2010). Therefore, if the phenotype of the ABIN1[D485N] mice is driven only by alterations in B cell function, it would be difficult to explain this solely by the loss of ABIN1-dependent recruitment of A20 to its substrates.

ABIN1 binds linear polyubiquitin chains, as well as K63-polyubiquitin chains (Fig. 1), and LUBAC, an E3 ligase which produces linear polyubiquitin specifically in vitro, appears to participate in at least one MyD88-dependent signaling pathway because the IL-1–stimulated activation of the canonical IKK complex was reported to be impaired in MEFs deficient in HOIL-IL, a component of LUBAC (Tokunaga et al., 2009). Therefore, the possibility that ABIN1 restricts signaling to NF-κB by binding to linear polyubiquitin chains as well as, or instead of, K63-pUb chains cannot be excluded at this stage. However, the activation of JNK by TNF was not inhibited in HOIL-IL−/− MEFs, implying that LUBAC is not required for the activation of TAK1. Moreover, the TAB2 component of the TAK1 complex binds to K63-pUb tetramers >100-fold more strongly than to linear ubiquitin tetramers (Kanayama et al., 2004; Kulathu et al., 2009; Sato et al., 2009). WT ABIN1 might therefore compete with the polyubiquitin-binding components of the TAK1 complex (TAB2–TAB3) for binding to the same K63-pUb chains/proteins, limiting the extent of activation of TAK1. We also observed in the present study that the endogenous polyubiquitylated IRAK1 could be communoprecipitated with WT ABIN1, but not with ABIN1[D485N], from the extracts of LPS-stimulated BMDM. Therefore, another possibility is that the interaction of WT ABIN1 with polyubiquitylated IRAK1 may hinder the ability of IRAK1 to interact with and activate TRAF6, reducing the TRAF6-mediated formation of K63-pUb chains and limiting the activation of TAK1. IRAK1 undergoes Lys63-linked polyubiquitylation in response to IL-1 (Windheim et al., 2008), but the possibility that it also undergoes linear polyubiquitylation has not been excluded.

Interestingly, while this study was in progress, human polymorphisms were identified in the gene encoding ABIN1 that predispose to lupus–like autoimmune diseases, psoriasis (Han et al., 2009; Nair et al., 2009), and vasculitis (He et al., 2010b). This suggests that ABIN1 also plays a key role in preventing autoimmunity in man. The ABIN1[D485N] mice may therefore be a good model for human autoimmune disease and could be used to assess the efficacy of drugs that target components of TLR–MyD88 signaling pathways. Drugs that increase the expression of ABIN1 could also have therapeutic value because the adenosivirus-mediated delivery of ABIN1 to the lung epithelium has been reported to reduce allergen-induced eosinophil infiltration of the lungs in a mouse model of allergen-induced asthma (El Bakkouri et al., 2005).

MATERIALS AND METHODS
Expression and purification of ABIN1 and NEMO and binding to polyubiquitin. Human ABIN1 and ABIN1[D472N] were cloned and expressed as fusion proteins with glutathione S-transferase (GST) at the N terminus and a His, tag at the C terminus. The expressed proteins were purified on nickel-nitritotriacetate agarose (Qiagen) followed by chromatography on glutathione-Sepharose (GE Healthcare). Human NEMO and NEMO[D311N] were expressed as GST fusions and purified on glutathione-Sepharose. 8 μg of these proteins were immobilized individually on a 10-μl packed volume of glutathione-Sepharose and incubated for 1 h at 21°C with 2 μg of K48-linked, K63-linked (Boston Biochem), or linear ubiquitin oligomers (Enzo Life Sciences) in 300 μl 25 mM HEPS, pH 7.5, 1 mM EGTA, 0.5% (vol/vol) Triton X-100, and 2 mM MgCl2. (Buffer A) plus 150 mM NaCl. The beads were washed five times with Buffer A plus 250 mM NaCl and once with Buffer A without Triton X-100.

Generation of ABIN1[D485N] knockin mice. The Asp485 to Asn knockin mutation was generated in exon 14 of the ABIN1 locus and, at the same time, LoxP sites were inserted in the introns adjacent to exons 14 and 16 by standard gene targeting methods (Fig. S1 A). The following primers were used for making the targeting vector (underlined regions indicate the presence of introduced restriction sites or LoxP sequences): 5′-GACCCGGCGCCAGACCTTTTTCATGCTAACAATACAGTTG-3′, 5′ arm sense (5′-AGGCCGCCAAGAGTGCCATACTCTGCGATATCAGGTAGTGAGTTGGATAGTTGTGG-3′), 3′ arm antisense (5′-AGGCCGCCATACCTCGTATACTCAGATATCAGGTAGTGAGTTGGATAGTTGTGG-3′), splice donor sense (5′-TGACCCGCCGCGCTGGAGCTAGTACCTGACCTAGT-3′), splice donor antisense (5′-TACCCGTTTAAATATACATGTCGTCGACGATCCCTGATCTCTTGCCTCCTCCAC-3′), and 3′ arm antisense (5′-ACTCGAGTCGACGATCCTCACCTTCTCGTCCCTCCAC-3′).

Primers used for screening and routine genotyping are as follows: P1 5′ LoxP sense (5′-ATGAAATTTCCTCCGACGATCAATCC-3′), P2 5′ LoxP antisense (5′-GAGGCGGTGATCTCGGAAATACC-3′), P3 mRNA sense (5′-TGTTGTTGAATGTTAGTTGTGG-3′), P4 mRNA antisense (5′-AGCTGCGCTGTGAAGATGAACG-3′), P5 3′ LoxP sense (5′-CCTCTCCATGGTCATGCC-3′), P6 3′ LoxP antisense (5′-AGCGTGACCTCGGAGCTGGATGTCGAGGG-3′), and P7 neo-ires sense (5′-CGTATTCACAAAGGATGCTAAGATGC-3′). All animals were maintained in specific pathogen-free conditions consistent with EU and UK regulations. All the work was performed under a UK Home Office project license that was awarded after recommendation by the University of Dundee Ethical Review Committee. The ABIN1[D485N] mice generated were on a 129Sv×C57BL/6 background and were backcrossed two generations to C57BL/6. MyD88−/− mice were provided by S. Akira (Osaka University, Osaka, Japan) and RIP2−/− mice were obtained from R.A. Flavell (Yale University School of Medicine, New Haven, CT).

Flow cytometry. Single-cell suspensions were made from thymus and spleen by gentle sieving and pipetting, whereas BM samples were prepared by flushing them out from tibia and femur with PBS. Peritoneal lavage was collected by injecting 10 ml PBS into the peritoneum. Erythrocytes were removed from spleen, BM, and peritoneal lavage by treatment with red blood cell lysis buffer (Sigma-Aldrich). For surface staining, cells were washed twice with ice-cold PBS, 2% (wt/vol) BSA, and 0.1% (wt/vol) sodium azide. 106 cells were blocked for 15 min with FcR antibody (purified CD16/32; 2.4G2; BD) and incubated for 30 min at 4°C with various antibodies conjugated to FITC, phycoerythrin (PE), PE-Cy5 (Tricolor), APC, or APC-Cy7 for multiple color fluorescence surface staining. For analysis of various cell types, cells were blocked for 15 min with FcR antibody (purified CD16/32; 2.4G2; BD) and incubated for 30 min at 4°C with various antibodies conjugated to FITC, phycoerythrin (PE), PE-Cy5 (Tricolor), APC, or APC-Cy7 for multiple color fluorescence surface staining. For analysis of various cell types, cells were blocked for 15 min with FcR antibody (purified CD16/32; 2.4G2; BD) and incubated for 30 min at 4°C with various antibodies conjugated to FITC, phycoerythrin (PE), PE-Cy5 (Tricolor), APC, or APC-Cy7 for multiple color fluorescence surface staining. For analysis of various cell types, cells were blocked for 15 min with FcR antibody (purified CD16/32; 2.4G2; BD) and incubated for 30 min at 4°C with various antibodies conjugated to FITC, phycoerythrin (PE), PE-Cy5 (Tricolor), APC, or APC-Cy7 for multiple color fluorescence surface staining.
cell extract protein was incubated for 2 h with 0.5 µg of the ABIN1 antibody.

Immunoblotting and immunoprecipitation. Cell extracts were prepared as previously described (Cheung et al., 2003), except that α-TAK1 was used to immunoprecipitate TAK1 instead of α-TAB1. In brief, TAK1 complexes were immunoprecipitated from 0.15 mg BMDC extract protein, using 0.45 µg anti-TAK1 antibody. The TAK1 activity in the immunoprecipitates was measured by the activation of MAPK kinase 6 and coupled to the activation of p38 MAPK. The active p38 MAPK generated in this step was quantified in a second assay by measuring its ability to incorporate [32P]-phosphate from γ-[32P]ATP into myelin basic protein.

Measurement of autoantibodies and immunoglobulin isotypes by ELISA. Blood was collected from mice by cardiac puncture, allowed to clot, and the serum was separated by centrifugation. Immunoglobulins (IgG, IgM) were measured by ELISA.

Multiplex cytokine assays. Cytokines in cell culture supernatants and mouse serum were measured using either Luminex-based Bio-Plex Mouse Cytokine 8-Plex, 6-Plex Panel, or 13 plex (Bio-Rad Laboratories) or multiplex cytokine assays (Millipore Corporation). Cytokines were measured by ELISA.

B cell and T cell proliferation. Untouched naive splenic B cells of >98% purity were isolated using a B cell isolation kit (Miltenyi Biotech) and 10^6 cells were cultured for 72 h in 96-well plates with LPS and LTA, R.848, anti-ιgM (Jackson ImmunoResearch Laboratories, Inc.), or anti-CD40 (Clone 3/23; BD). Splenic and LN CD4^+ T cells (>95% pure) were isolated using a CD4^+ T cell isolation kit (Miltenyi Biotech) and stimulated for 72 h with plate-bound anti-CD3 alone (clone 145-2C11; BD) or with anti-CD3 plus anti-CD28 (37.5:1; BD). The samples were pulsed for 16 h with 0.5 µCi [3H]thymidine before harvesting and measuring [3H]thymidine incorporation.

Generation of BMDM and BMDC. BMDM and BMDC were obtained by differentiating femur and tibia BM using recombinant mouse M-CSF (R&D Systems) and GM-CSF (R&D Systems), respectively. After 12–24 h of replating, cells were stimulated with LPS (Escherichia coli strain O5:B5; Enzo Life Sciences), LTA from Staphylococcus aureus, Pam3CSK4, CpG DNA (ODN 1826), or R.848 (TLR7 agonist), all obtained from InvivoGen. For immunoprecipitation and immunoblotting, cell extracts were prepared as previously described (Windheim et al., 2008).

Immunoblotting and immunoprecipitation. 10 µg of cell extract protein was subjected to SDS-PAGE and immunoblotting at high sensitivity. After washing the membrane over end for 60 min with 15 µl of protein G–Agarose beads, the agarose was collected and washed six times with cell lysis buffer plus 0.5 M NaCl. Immunoprecipitates were denatured in SDS, subjected to SDS-PAGE, and immunoblotted.

Antibodies for immunoblotting and immunoprecipitation. Sheep antibodies against mouse ABIN1 (sheep number S345C, third bleed) and human TAK1 (sheep number S828A, first bleed) were generated in the MRC Protein Phosphorylation Unit by J. Hastie (University of Dundee, Dundee, UK). Antibodies that recognize IKK-α/β phosphorylated at Ser180/Ser181 or Ser176/180, p105/NIκB phosphorylated at Ser933, p38 MAPK phosphorylated at the TGY motif, ERK1/ERK2 phosphorylated at the TGY motif, βc/6ε phosphorylated at Ser32/36, and antibodies that recognize the deubiquitylase A20, all forms of p38 MAPK, ERK1/ERK2, βc/6ε, Caspase8, and Caspase3 were obtained from Cell Signaling Technology. JNK phosphorylated at the TYP motif was from Invitrogen. Mouse monoclonal antibodies against GAPDH (Abcam) and ubiquitin (P0D1; Santa Cruz Biotechnology, Inc.), anti-IRAK1 (Santa Cruz Biotechnology, Inc.), and rabbit-, mouse-, and sheep-specific secondary antibodies conjugated to hors eradish peroxidase (Thermo Fisher Scientific) were purchased from the suppliers in parentheses.

Measurement of TAK1 activity. This was performed as described previously (Cheung et al., 2003), except that α-TAK1 was used to immunoprecipitate TAK1 instead of α-TAB1. In brief, TAK1 complexes were immunoprecipitated from 0.15 mg BMDC extract protein, using 0.45 µg anti-TAK1 antibody. The TAK1 activity in the immunoprecipitates was measured by the activation of MAPK kinase 6 and coupled to the activation of p38 MAPK. The active p38 MAPK generated in this step was quantified in a second assay by measuring its ability to incorporate [32P]-phosphate from γ-[32P]ATP into myelin basic protein.

Statistical analysis. Statistical significance was calculated either using the two-tailed Student's t test or the Mann-Whitney U test using Prism software (GraphPad Software). Further details are provided in the relevant figure legends.

Online supplemental material. Fig. S1 shows design of the targeting vector and the expression pattern of ABIN1 protein in various cells from ABIN1[D485N] mice. Fig. S2 shows immune phenotyping of ABIN1[D485N] mice. Fig. S3 shows no alteration in T cell receptor signaling in ABIN1 knockin mice. Fig. S4 shows enhanced activation of LPS signaling in B cells and also enhanced activation of TLR-MyD88 and NOD1-RIPI signaling pathways in BMDM from ABIN1[D485N] mice. Fig. S5 shows rescue of the T cell phenotype observed in ABIN1[D485N]/MyD88^−/− mice. Fig. S6 shows enhanced activated of TLR signaling in BMDC from ABIN1[D485N]/MyD88^−/− mice. Fig. S6 shows enhanced activated of TLR signaling in BMDC from ABIN1[D485N]/MyD88^−/− mice. Fig. S6 shows enhanced activated of TLR signaling in BMDC from ABIN1[D485N]/MyD88^−/− mice. Fig. S6 shows enhanced activated of TLR signaling in BMDC from ABIN1[D485N]/MyD88^−/− mice. Fig. S7 shows TNF and IL-1 signaling in embryonic fibroblasts from ABIN1[D485N] mice. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20101777/DC1.
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