The ubiquitin-editing enzyme A20 controls NK cell homeostasis through regulation of mTOR activity and TNF

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The ubiquitin-editing enzyme A20 is a well-known regulator of immune cell function and homeostasis. In addition, A20 protects cells from death in an ill-defined manner. While most studies focus on its role in the TNF-receptor complex, we here identify a novel component in the A20-mediated decision between life and death. Loss of A20 in NK cells led to spontaneous NK cell death and severe NK cell lymphopenia. The few remaining NK cells showed an immature, hyperactivated phenotype, hallmarked by the basal release of cytokines and cytotoxic molecules. NK-A20−/− cells were hypersensitive to TNF-induced cell death and could be rescued, at least partially, by a combined deficiency with TNF. Unexpectedly, rapamycin, a well-established inhibitor of mTOR, also strongly protected NK-A20−/− death and could be rescued, at least partially, by a combined deficiency with TNF. This study therefore maps A20 as a crucial regulator of mTOR signaling and underscores the need for a tightly balanced mTOR pathway in NK cell homeostasis.

**BRIEF DEFINITIVE REPORT**

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## Introduction

Natural killer (NK) cells are the cytotoxic members of the heterogeneous population of innate lymphoid cells (ILCs; **Vivier et al., 2018**). NK cells kill target cells via the binding of death receptors or by the release of lytic granules that contain granzymes and perforin. They also regulate the function of other immune cells by producing chemokines and cytokines such as TNF and IFNγ (**Vivier et al., 2008**). Under normal conditions, their activation is inhibited by ligands expressed on healthy cells that engage germline-encoded inhibitory receptors on the NK cells. Viral infection (**Waggoner et al., 2016**), malignant transformation (**Vivier et al., 2012**), or cellular stress (**Raulet and Guerra, 2009**) can lead to up-regulation of ligands that are recognized by a vast array of activating receptors. The relative balance of inhibitory and activating signals eventually determines the activity of the NK cell. Several signaling pathways have been identified to play a crucial role in NK cell functioning. Recently, the mechanistic target of rapamycin (mTOR) pathway was shown to be a hallmark of NK activity (**Marçais et al., 2014, 2017**). Although NK cell activation has been studied thoroughly, relatively little is known about how activated NK cells are switched off after termination of an inflammatory response.

The NF-κB family of transcription factors plays a key role in inflammatory responses triggered by a plethora of signaling receptors. NF-κB dimers induce expression not only of a large proinflammatory gene program, but also of their own negative regulators, such as inhibitor of κB (IκB) or A20 (encoded by the gene TNFAIP3 induced protein 3 (**Tnfaip3**; **Renner and Schmitz, 2009**). Absence of A20 leads to prolonged NF-κB activation and elevated production of inflammatory cytokines (**Catrysse et al., 2014**). A20-deficient mice show widespread tissue inflammation and perinatal death (**Lee et al., 2000**). Also in humans, polymorphisms in the **Tnfaip3** gene are...
A20-deficient NK cells are hyperactive

NK cells differentiate through a series of maturation stages, which can be traced in vivo by the sequential acquisition and loss of specific cell surface receptors. Immature NK cells are defined by the presence of CD27 and absence of CD11b (CD11b<sup>lo</sup>), while fully mature NK cells gain CD11b and lose CD27 expression (CD27<sup>hi</sup>–; Kim et al., 2002; Hayakawa and Smyth, 2006; Chiossone et al., 2009; Fig. 2 A). Although NK cell numbers were severely decreased in each maturation stage (Fig. 2 A, lower panel), the relative proportions of the different subtypes changed dramatically, and residual NK cells in the spleen of NK-A20<sup>−/−</sup> mice displayed a marked shift toward the immature phenotype (Fig. 2 A, contour plot and middle panel). Furthermore, A20-deficient NK cells showed a highly proliferative phenotype as assessed by Ki-67 expression (Fig. 2 B, upper panel). This enhanced proliferation was not a direct consequence of having lower numbers of NK cells in an attempt to fill up the empty niche, as A20-deficient NK cells in mixed BM chimera mice reconstituted with WT and NK-A20<sup>−/−</sup> BM also showed increased Ki-67 expression (Fig. 2 B, lower panel).

We then performed functional studies and tested intracellular IFNγ production and degranulation, as measured by cell-surface CD107a expression, a lysosomal marker that translocates to the cell surface upon degranulation (Alter et al., 2004). Even without prior stimulation, NK-A20<sup>−/−</sup> mice displayed a hyperactive phenotype as revealed by the enhanced proportion of CD107a<sup>hi</sup>– and IFNγ<sup>−/+</sup> NK-A20<sup>−/−</sup> cells in steady state (Fig. 2 C). Ex vivo restimulation with plate-bound antibodies against NK1.1 or with the cytokines IL-12/IL-18 or IL-12/IL-2 triggered a higher degree of degranulation in the absence of A20. The proportion of IFNγ<sup>−/−</sup> cells was not altered, but we noticed that IFNγ expression levels on a per-cell basis were strongly elevated in NK-A20<sup>−/−</sup> cells (Fig. 2 D). Finally, we also assessed NF-κB activation, using phosphorylation of Ser536 on p65 (P-p65) as readout. Both in steady state and upon TNF activation, levels of P-p65 were markedly increased in the absence of A20 (Fig. 2 D), as expected from A20’s known NF-κB inhibitory function.

In summary, these data show that the few remaining NK cells in the NK-A20<sup>−/−</sup> mice showed a relatively immature phenotype but were hyperactive, as revealed by increased proliferation, increased cytokine production, increased degranulation, and increased signs of NF-κB activation, both in steady state and upon ex vivo restimulation.

A20 is indispensable for NK cell survival

Our results thus far showed that NK-A20<sup>−/−</sup> mice have severe NK cell lymphopenia. To assess whether A20 might affect NK cell survival, we first examined the viability of freshly isolated splenocytes by staining with annexin V and a viability marker. We consistently measured decreased viability of A20-deficient NK cells, accompanied by slightly activated caspase 3 levels (Fig. S2, A and B). To follow the fate of NK-A20<sup>−/−</sup> cells over time, we generated a tamoxifen-inducible A20 knockout mouse by crossing LoxP-flanked A20 mice to the ERT2-Cre<sup>lox/lox</sup> mouse line. As full A20 deletion is lethal (Lee et al., 2000), we enriched NK cells from the spleens of these mice (CD45.2<sup>Lo</sup> A20<sup>lox/lox</sup> × ERT2-Cre<sup>lox/lox</sup> or ERT2-Cre<sup>−/−</sup>) and mixed them with enriched NK cells from
Figure 1. Genetic ablation of A20 in Nkp46+ cells leads to severe NK cell lymphopenia. (A) Graphical representation and validation by RT-qPCR of the NK-A20 mouse model. Bar graphs represent A20 RNA expression as measured in sorted splenic lymphocyte populations. Expression was normalized to the indicated housekeeping genes. (B and C) Flow cytometry analysis to assess NK cell percentages in the indicated organs and absolute numbers of splenic lymphocyte populations in NK-A20 mice. (D) Graphical representation of NK-A20 mixed BM chimeric setup. (E–G) FACS plots and bar graphs represent percentages or ratios of CD45.1.2+ and CD45.2+ B cells or NK cells as retrieved from the spleens or indicated organs of mixed BM chimeras. For all figures, bar graphs represent mean ± SD, and dots represent individual mice. Data are representative of at least two independent experiments, with the exception of A, middle panel. Unpaired Student’s t tests: *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.
Figure 2. **Loss of A20 renders NK cells relatively immature and highly hyperactive.** (A–D) Flow cytometry analysis of splenic NK cells isolated from NK-A20 mice. (A) FACS plots and bar graphs represent percentages and absolute numbers of NK cell subsets, determined by CD11b/CD27 expression. CD11blo, CD11bloCD27hi, immature; DP, CD11bhiCD27hi, double positive (DP) or intermediate; CD27lo, CD11bloCD27hi, mature. (B) Ki-67 expression as measured in splenic NK cells from NK-A20 mice (top) or mixed BM chimeric mice (bottom). (C) Bar graphs depict percentages of CD107a+ or IFNγ+ NK cells and IFNγ geometrical mean fluorescence intensity (MFI) within IFNγ+ NK cell population. (D) P-p65 MFI on gated splenic NK cells as measured by phosphoflow immediately after isolation or upon TNF stimulation. For all figures, bar graphs represent mean ± SD, and dots represent individual mice. Data are representative of at least two independent experiments. Unpaired Student’s t test: **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.
CD45.1 congenic control mice (Fig. 3 A). This mix was adoptively transferred to CD45.1.2 congenic acceptor mice that were subsequently injected i.p. with tamoxifen for 5 d consecutively. Acceptor mice were bled on days 1, 4, 7, and 10 to determine the ratio of CD45.2 to CD45.1 donor NK cells, which was compared to the ratio before injection (Fig. 3 A, flow panels and survival curve). This analysis showed that upon tamoxifen injection, ERT2-Cre+ NK cells lost A20 expression from day 2 onwards (Fig. 3 A, bar graph) and were rapidly depleted, unequivocally demonstrating the essential role for A20 in NK cell survival (Fig. 3 A).

NK cell homeostasis depends on IL-15 signaling, which drives expression of the antiapoptotic protein Mcl-1 and inhibits expression of proapoptotic Bim (Huntington et al., 2007; Sathe et al., 2014). Under IL-15-depriving conditions, this delicate balance of pro- and antiapoptotic Bcl-2 family members is disturbed: Bim and Noxa levels rise, while Mcl-1 levels go down. Ectopic Bcl-2 expression restores this balance and can inhibit NK cell apoptosis (Cooper et al., 2002; Minagawa et al., 2002). To determine potential changes in expression of Bcl-2 family members in the absence of A20, we analyzed their expression specifically in the CD11b+ subset, to avoid any potential bias due to the skewed maturation of A20−/− NK cells. While levels of Bcl-2, Bcl-xL, Mcl-1, XIAP, cIAP1, and Bim appeared largely unaltered, we found a marked increase in the levels of the proapoptotic molecule Noxa in A20−/− NK cells (Fig. S2 C). To determine if Bcl-2 can restore NK cell survival in the absence of A20, NK-A20−/− mice were crossed with H2K-bcl-2 transgenic mice expressing human Bcl-2 in all hematopoietic cells including NK cells (Domen et al., 1998). Bcl-2Tg/+ mice displayed splenomegaly, and numbers of splenic NK cells were elevated compared with littermate control mice, as described for other hematopoietic cells (Domen et al., 2000). Notably, the NK cell fold expansion induced by ectopic Bcl-2 expression was significantly higher in A20−/− compared with A20+/+ NK cells (7.6- vs. 1.5-fold; Fig. 3 B, lower panel), suggesting that A20−/− NK cells die, at least partially, through an apoptotic pathway dependent on the intrinsic apoptotic pathway. The phosphorylation status of STAT5 was not significantly different between NK-A20−/− and littermate NK cells either in basal conditions or upon stimulation with IL-15, indicating that signaling immediately downstream of the IL-15 receptor was not affected by loss of A20 (Fig. 4 A). Along these lines, the levels of CD22 (IL-2/IL-15 receptor β chain) were unaltered in NK-A20−/− cells (data not shown). Recent data implemented a role for the metabolic checkpoint kinase mTOR in IL-15-dependent signaling (Marçais et al., 2014). Notably, in T cells, mTOR activity was enhanced upon loss of A20 (Matsuzawa et al., 2015). Also, NK-A20−/− cells showed strongly enhanced mTORC1 activity.

A20-deficient NK cells display elevated mTOR activation

As NK cell homeostasis is intricately associated with IL-15 and STAT5 signaling (Cooper et al., 2002; Vosshenrich et al., 2005; Eckelhart et al., 2011), we assessed whether A20 might modulate this pathway. The phosphorylation status of STAT5 was not different between NK-A20−/− and littermate NK cells either in basal conditions or upon stimulation with IL-15, indicating that signaling immediately downstream of the IL-15 receptor was not affected by loss of A20 (Fig. 4 A). Along these lines, the levels of CD22 (IL-2/IL-15 receptor β chain) were unaltered in NK-A20−/− cells (data not shown). Recent data implemented a role for the metabolic checkpoint kinase mTOR in IL-15-dependent signaling (Marçais et al., 2014). Notably, in T cells, mTOR activity was enhanced upon loss of A20 (Matsuzawa et al., 2015). Also, NK-A20−/− cells showed strongly enhanced mTORC1 activity.
A20mZnF7/mZnF7 mice suffer from massive hyperinflammation tamoxifen-induced cell death of ERT2-Cre+ A20-deficient NK cells, as determined by annexin V and live/dead staining on flow cytometry. Bar graphs show AUC for indicated mTOR substrates per genotype. For readability, significant differences are shown only per genotype but were also evident between genotypes (****, P < 0.0001 per dose).

Figure 3. A20 is indispensable for NK cell survival. (A) Graphical representation of adoptive cell transfer setup. FACS plots show the frequency of CD45.1 and CD45.2 donor NK cells, and survival graph shows CD45.2/CD45.1 donor NK cell ratios compared with day 0 as measured over time in the bar graph. FACS plots depict the proportion of NK cells per genotype. (B) Freshly isolated NK-A20 splenocytes were subjected to a dose range of mTNF over time. Survival curves (upper graph) represent NK cell viability as assessed by annexin V and live/dead staining on flow cytometry. Bar graphs (lower panel) show AUC for indicated mTNF doses per genotype.

To assess whether this enhanced mTOR activity was contributing to the observed NK-A20−/− cell death, we injected CD45.1 acceptor mice for 6 d consecutively with rapamycin, an inhibitor of mTORC1, to see if we could restore tamoxifen-induced cell death of ERT2-Cre+ A20-deficient NK cells. As can be seen in Fig. 4 D (middle panel), we obtained a clear rescue of A20−/− NK cells by damped the exacerbated mTOR signaling by rapamycin, classifying mTOR as a bona fide target in the A20-mediated decision between life and death.

To test whether the TNF and mTOR pathways act independently or belong to one and the same pathway, we concomitantly blocked both TNF and mTOR during the tamoxifen-induced deletion process of A20 and assessed whether the two inhibitors worked synergistically. As can be noted from Fig. 4 D (lower panel), each inhibitor individually rescued part of the NK cells, but together they led to a complete restoration of NK cell death induced by loss of A20. This suggests that both TNF hypersensitivity and exacerbated mTOR signaling contribute separately to NK cell death in conditions of A20 deficiency.

A20 comprises a deubiquitinase domain as well as ubiquitin-binding domains. Our department recently generated a ubiquitous zinc finger 7 (ZnF7) knock-in mouse model for A20, in which cysteine residues 764 and 767 in ZnF7 domain were mutated (referred to as A20mZnF7) to abrogate binding of A20 to linear ubiquitin chains in all cells (Polypkratis et al., 2019). A20mZnF7/mZnF7 mice suffer from massive hyperinflammation and develop arthritis, similar to mice deficient for A20 in myeloid cells (Polypkratis et al., 2019). To elucidate how A20 mediates its homeostatic role in NK cells, we immunophenotyped the splenic NK cell compartment in the A20mZnF7/mZnF7 mice and found that impairment of A20 linear ubiquitin binding causes massive cell death in NK cells (Fig. 5 A). Similar to what was shown in NK-A20−/− mice, mTOR activity was highly elevated in A20mZnF7/mZnF7 cells (Fig. 5 B), while intracellular TNF levels in A20mZnF7/mZnF7 cells did not differ (Fig. 5 C). However, at the serum level, we found a massive increase in TNF in the A20mZnF7/mZnF7 mice (Fig. 5 D).

In summary, we unraveled a cell-intrinsic role for A20 in controlling NK cell survival by acting on the TNFRI and mTOR complexes. Constitutive loss of A20 in NK cells led to severe and widespread NK cell lymphopenia due to NK cell death. Remaining NK cells were hyperactive and hyperproliferative, produced more proinflammatory cytokines, and had increased cytotoxic properties. Despite the well-established anti-apoptotic role of A20 (Lee et al., 2000), NK cells are, to our knowledge, the first cell type that die in a spontaneous manner upon loss of A20. B cells and dendritic cells show strongly enhanced survival, due to the up-regulation of Bcl-2 and Bcl-XL, and are even more resistant to Fas-induced cell death (Tavares et al., 2010; Kool et al., 2011). Strikingly, despite strongly enhanced activation of the NF-κB pathway in NK-A20−/− cells (as monitored by P-p65 levels), antiapoptotic NF-κB–dependent genes were not induced, showing complex regulation of NF-κB transcriptional activity in NK cells. In T cell–specific conditional A20−/− mice, frequency and numbers of CD4+CD8+ double-positive thymocytes, CD8− or CD4+ single-positive thymocytes, and peripheral T cells were normal (Onizawa et al., 2015). Only upon TCR triggering could marked reductions in T cell numbers be observed, suggesting that A20 was needed to protect activated T cells from dying (Onizawa et al., 2015). Similarly, in hepatocytes and enterocytes, loss of A20 did not lead to spontaneous cell death, but rather caused hypersensitivity to sublethal doses of TNF (Vereecke et al., 2010; Catrysse et al., 2016). In NKT cells, A20 controlled survival of NKT1 and NKT2 to some extent, and not NKT17 subsets, but never led to the dramatic cell loss observed for NK- A20−/− cells (Drennan et al., 2016). Loss of A20-deficient NKT cells could be prevented by compound deficiency with MALT1, a proteolytic enzyme in the Carma-Bcl10–MALT1 complex downstream of the TCR and a well-established target of A20. However, this pathway appeared not to be involved in A20-mediated cell death of NK cells (data not shown). Part of the mechanism leading to spontaneous loss of NK cells in the absence of A20 was explained by the hyperactivated state of A20−/− NK cells, which led to increased proinflammatory TNF production, either direct or indirect. Due to the loss of A20 and
**Figure 4.** **A20 restricts the metabolic activity of NK cells.** (A and B) Flow cytometry analysis of splenic NK cells isolated from NK-A20 mice. (A) Histograms show steady state phosphorylation (p-) of indicated proteins in CD11b\(^{lo}\) NK cells measured by phosphoflow. Bar graphs depict mean fluorescence intensity (MFI) of phosphorylated proteins in steady state or upon IL-15 stimulation. FMO, fluorescence minus one. (B) Histograms show 2-NBDG uptake and CD71 and CD98 expression as measured on steady state CD11b\(^{lo}\) NK cells. Bar graphs depict MFI of the indicated parameters in NK cell subsets. DP, double positive. (C) Graphical representation of OP-Puro incorporation in NK-A20 BM chimeras. Bar graphs represent OP-Puro fluorescence as measured by flow cytometry in splenic donor-derived CD45.1.2 (WT) or CD45.2 lymphocyte populations. (D) Graphical representation of adoptive cell transfer setup. Bar graphs depict the
concomitant hypersensitivity to TNF-induced killing, this caused NK cell death. Still, as was obvious from the compound deficient NK-A20−/−×TNF−/− mice, NK cell numbers could not be restored to WT levels, indicating that additional mechanisms were needed to explain the A20-mediated protection against cell death.

We turned our attention toward the mTOR signaling cascade, a pathway that was recently established as a critical regulator of NK cell homeostasis and considered a hallmark for reactive NK cells (Marçais et al., 2014, 2017). Loss of A20 dramatically increased mTOR signaling output, as monitored by enhanced downstream phosphorylation of 4EBP1 and S6, enhanced glucose uptake, and enhanced translation. This occurred independently of IL-15 receptor signaling, as both IL-15 receptor levels and signaling steps immediately downstream of the IL-15 receptor were unaffected. Rather, we propose that A20 could directly target either mTOR or any of its regulators in the mTOR signaling complex by ubiquitination, as suggested before for T cells (Matsuzawa et al., 2015). It has been shown that increased ubiquitination of mTOR and/or proteins within the mTOR signaling complex are associated with restrained mTOR activity, in an as yet ill-defined manner (Ivanov and Roy, 2013). Indeed, a newly generated A20mZnF7 knock-in mouse model confirmed that ZnF7 is absolutely essential for A20 to restrict mTOR activity and preserve NK cell homeostasis, implying a potential role for A20 in stabilization of M1 ubiquitin chains on mTOR or associated proteins.

We can only speculate how increased mTOR signaling causes NK cell death. Activation of mTOR has been associated before with increased NK cell reactivity, resulting in enhanced calcium influxes upon triggering of NK activating receptors (Marçais et al., 2017). Furthermore, increased translation is linked with increased production of oxygen radicals (Han et al., 2013). While not formally proven, it is not difficult to conceive that both calcium overload and ROS production could trigger mitochondrial pathways of cell death. Intriguingly, we observed a robust increase in the expression level of the proapoptotic molecule Noxa in NK-A20−/− cells and found that the cells could be partially rescued by overexpression of Bcl-2, suggesting at least a partial contribution of a mitochondrial pathway of death. Finally, high mTOR activation is also known to interfere with cytoprotective pathways of autophagy. Preliminary data showed that deficiency of Atg16L1, a crucial component of autophagosome formation (Fujita et al., 2008; Adolph et al., 2013), also causes NK cell loss in a dose-dependent manner (data not shown). These data confirm that a tight balance of mTOR signaling and autophagy is crucial for proper NK cell homeostasis (O’Sullivan et al., 2016).

In conclusion, here we established a critical role for A20 in restricting mTOR activity in NK cells, adding a novel player in the decision of A20 between life and death. We postulate that NK cells, being exquisitely sensitive to balances between inhibitory and activating signals, are particularly vulnerable to any dysregulation of these balances and therefore prone to death when a crucial regulator such as A20 is lost.

Materials and methods

Mice

The Ncr1Cre/− mouse line was originally described in Narnimancinelli et al. (2011); A20fl/fl mice in Vereecke et al. (2010); RIPK3−/− in Newton et al. (2004); H2K-bcl-2g/− in Domen et al. (1998); TNF−/− mice in Pasparakis et al. (1996); and A20mZnF7 mice in Polykrati et al. (2019). The ERT2-Crebg/−, CD45.1, and CD45.1.2 mice are provided by the Jackson Laboratory. The following mouse lines were bred and housed under specific pathogen-free conditions at the VIB-Ghent University Center for Inflammation Research: Ncr1−/−×A20flbg/−, Ncr1Cre/−×A20flbg/− or A20flbg/− (NK-A20−/−, NK-A20−/−, or NK-A20−/−); NK-A20−/−×RIPK3−/−, NK-A20−/−×TNF−/−, A20flbg/−×ERT2-Crebg/−, CD45.1, CD45.1.2, and A20mZnF7. The NK-A20−/−×H2K-bcl-2g/− mouse line was housed at the Campus Proeftuinstraat Ghent University site. All transgenic mouse lines have been generated on a C57Bl/6 background (Ncr1Cre/−, A20flbg/−, ERT2-Crebg/−), and A20mZnF7 or were backcrossed for >10 generations to C57Bl/6 (H2K-bcl-2g/−, RIPK3−/−, TNF−/− transgenic mice). All experiments with the NK-A20 mice were performed with NK-A20−/−/− animals as littermate controls. In the case of adoptive transfers or BM chimeras, CD45.1/2 congenically marked animals were included as WT counterparts. All animal experiments were performed in accordance with institutional guidelines for animal care of the VIB site Ghent/Ghent University Faculty of Sciences and in accordance with ethical committee EC2014_043. Litters with mice of both sexes at 6–17 wk of age were used for experiments.

Generation of BM chimeras

CD45.1 recipient mice received two rounds of sublethal irradiation (400 cG) with a time interval of 4 h. At least 4 h after the second irradiation cycle, recipient mice were injected intravenously with a 50:50 mixture of BM cells (either 2 × 106 or 4 × 106) derived from CD45.1 and NK-A20−/−, NK-A20−/−, or NK-A20−/− donors. Mice were used for experiments ≥8 wk after reconstitution.

Preparation of single-cell suspensions

For all figures except Fig. S1, spleens and mediastinal lymph nodes were smashed over a 70-µM filter, and red blood cells were removed from the spleen by osmotic lysis. For Fig. S1, an enzymatic digestion protocol was followed that ensures proper isolation of the myeloid compartment. Spleens were first minced, digested for 30 min in RPMI 1640 (Gibco) containing Liberase TM (0.02 mg/ml; Roche) and DNase I (10 U/ml; Roche) in a shaking water bath at 37°C, filtered through a 70-µM filter, and subjected to osmotic lysis. For BM, single-cell suspensions were subjected to osmotic lysis. For BM, single-cell suspensions were subjected to osmotic lysis.
were obtained by crushing the tibia and femur, followed by lysis of red blood cells. In the case of lung and liver, organs were collected from PBS-perfused mice, minced, and digested at 37°C for 20–45 min in RPMI 1640 (Gibco) containing DNase I (10 U/ml; Roche) and collagenase A (1 mg/ml; Sigma-Aldrich). Lungs required a single GentleMACS dissociation step (after digestion), while livers were subjected to two rounds of GentleMACS dissociation (before and after digestion). Red blood cells were removed by osmotic lysis, and an additional Percoll-gradient centrifugation step was performed for the liver. To isolate cells from the lamina propria, the small intestine was dissected, Peyer’s patches were removed, and the intestine was opened longitudinally and then minced using scissors. Epithelial cells were removed by serial washes at 37°C with HBSS without Ca2+.  

Figure 5. A20 controls NK cell survival and the mTOR pathway through its ZnF7 domain. (A–C) Flow cytometry analysis of steady state splenic NK cells isolated from indicated mouse lines. (A) Bar graphs depict NK cell absolute numbers. (B) Histograms and bar graphs depict MFI of the indicated phosphorylated proteins as determined by phospho-flow. (C) Bar graphs show TNF MFI in total NK cells (no prior stimulation). All bar graphs represent mean ± SD, and dots show individual mice. All data are representative of at least two independent experiments, and statistical analysis was performed with a one-way ANOVA (ad hoc Tukey’s multiple comparisons test). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. (D) Model depicting the possible role of A20 in NK cells. Upper panel: Through its Zn fingers, A20 binds and stabilizes M1-linked ubiquitin chains in the TNF receptor complex. As such, A20 competes with NF-κB essential modulator (NEMO) for binding to key components in the receptor disc and thereby interferes with NF-κB activation. At the same time, by stabilizing M1 ubiquitin chains, A20 also prevents the deubiquitinase cylindromatosis (CYLD) from degrading linear ubiquitin chains. This precludes formation of the proapoptotic complex II. Lower panel: Loss of A20 leads to strongly enhanced NF-κB activation and release of cytokines, such as TNF, in either a direct or indirect manner. Due to the enhanced sensitivity to TNF-mediated killing in the absence of A20, NK cells die. Additionally, A20 also restricts mTOR activity in NK cells, preventing spontaneous hyperactivation of this pathway. Loss of A20 leads to mTOR overactivation and mTOR-dependent NK cell death, which could be due to enhanced translation and increased production of ROS, to a block in autophagy and/or other signaling pathways. LUBAC, linear ubiquitination assembly complex.
and Mg²⁺ (Gibco) containing 10% FCS (Bodinco) and 2 mM EDTA (Lanza). The lamina propria was subsequently digested at 37°C for 15–20 min in RPMI 1640 supplemented with 10% FCS, collagenase A (0.24 mg/ml; Sigma-Aldrich) and DNase I (50 U/ml; Roche), followed by Percoll-gradient centrifugation. Tail vein blood was collected in Microvette 200 K3E tubes (Sarstedt), and red blood cells were cleared by several rounds of osmotic lysis. Serum samples were obtained from femoral artery (terminal) bleeding. Blood was collected in Eppendorf tubes and stored at room temperature for ≥2–3 h. Samples were then centrifuged for 10 min at 5,000 rpm, and supernatant was collected and stored at −20°C.

Flow cytometry and gating strategy

Single-cell suspensions were obtained as described and stained with monoclonal antibodies labeled with fluorochromes or biotin, recognizing the following markers: activated caspase 3 (C92), CD3 (17A2 or 145-2c11), CD11b (M1/70), CD11c (N418), CD16/32 (2.4G2), CD19 (1D3), CD27 (LG.7F9 or LG.3A10), CD45 (30-F11), CD5 (B56), NK1.1 (PK136), NKp46 (29A1.4), P-4EBP1 (236B4), (X54-5/7.1), CD71 (R17217), CD98 (RL388), CD107a (1DB4), CD122 (2.4G2), CD19 (1D3), CD27 (LG.7F9 or LG.3A10), CD45 (30-F11), CD5 (B56), NK1.1 (PK136), NKp46 (29A1.4), P-4EBP1 (236B4), CD45R0 (5B6), NK1.1 (PK136), NKp46 (29A1.4), P-4EBP1 (236B4), P-AKT (M89-61), PDCDA1 (JF05-1C2.4.1), P-p65 (93H1), P-S6K (DS7.2.2E), P-STAT5 (47), Rorγt (AKF5S-9), SiglecF (E50-2440), TCRβ (H57-597), TCRγδ (eBioGL3), and TNF (MP6-XT22). Intracellular stainings for TNF and OP-Puro were used for intracellular stainings of phosphorylated proteins. Intracellular IFNγ, caspase 3, and Ki-67 stainings were performed with Cytofix/Cytoperm (BD) and Perm/Wash (BD). Cell viability was measured using annexin V (BD) and/or a fixable viability dye (eBioscience). Flow cytometry was performed on a FACS Fortessa 4 or 5 laser (BD Biosciences), and data were analyzed using FlowJo.

The following gating strategy was applied for NK-A20 immunophenotyping: B cells (live, CD3CD19+), T cells (live, CD19NK1.1CD3+), CD4 T cells (live, CD19NK1.1CD3+TCRγCD4+), CD8 T cells (live, CD19NK1.1CD3+TCRγδCD8+), macrophages (live, CD19NK1.1CD3+FCG4/80CD64+autofluorescent), dendritic cells (live, CD19NK1.1CD3−not macrophages) CD11c MHCIIm+, NK cells (live, CD19CD3NK1.1+), NK T cells (live, CD19CD3NK1.1+), neutrophils (live, CD19NK1.1CD3−not basophils) CD11bLy6G+, eosinophils (live, CD19NK1.1CD3−not basophils or neutrophils) CD11cSiglecF−SSc, Ly6Chigh monocytes (live, CD19NK1.1CD3−not basophils, neutrophils, or eosinophils) CD11bLy6C+, plasmacytoid dendritic cells (live, CD19NK1.1CD3−not basophils, neutrophils, eosinophils, or Ly6Chigh monocytes) Ly6C−PDCA1+, and γδ T cells (live, CD19NK1.1CD3+TCRγδ+). The following standard gating strategy was used for NK cells throughout the article: live, CD19CD3NK1.1CD122CD49b+. In FACS plots showing NK cells, numbers depict frequencies of NK cells of total live cells. Depending on antibody availability, CD122 was sometimes replaced by NKp46, and CD19 was not always added. For phosphoflow, the following NK cell gating was applied: live, CD3NK1.1NKp46+, or live, CD3NK1.1CD49b+. For some stainings, fluorescence minus one was used as a control.

Cell sorting and gating strategy

Leukocytes were obtained from the spleen as described and, before sorting, enriched for NK cells with the MagniSort NK cell enrichment kit (eBioscience) or with an in-house optimized enrichment procedure using MACS columns (negative selection with stainings against CD5, CD24, TCRβ, CD19, F4/80, CD3, Ter119, and Ly6G). For sorting T, B, and NK cells, the obtained single-cell suspension was divided in two fractions. One fraction was immediately stained for T and B cells, and the second fraction was used for NK enrichment before staining. Up to 40,000 cells (T, B, and NK cells) were sorted directly in RLT Plus Buffer (RNeasy plus micro kit; Qiagen) supplemented with β-mercaptoethanol (10 µl/ml). Samples were snap frozen and stored at −80°C until further processing. Cell sorting was performed on a FACS AriaII and III (BD Biosciences). The following gating strategies were applied: B cells (live, CD19+), T cells (live, CD19NK1.1CD3+), and NK cells (live, CD3TCRβCD19+CD22NK1.1CD49b+, or live, CD3CD19TCRβNK1.1NKp46+).

RNA extraction and quantitative RT-PCR (RT-qPCR)

RNA from sorted cells was obtained using the RNeasy Plus Micro Kit (Qiagen) according to the manufacturer’s instructions. cDNA was generated with the SensiFAST cDNA Synthesis Kit (Bioline) or the iScript cDNA synthesis kit (Bio-Rad Laboratories) or amplified with the Ovation PicoSL WTA System V2 kit (NuGEN) following the manufacturer’s guidelines. Upon amplification, pollutants were removed using the MinElute PCR purification kit (Qiagen) before SybrGreen-based RT-qPCR with the SensiFast SYBR No-ROX kit (Bioline) on a LightCycler 480 (Roche). The following mouse primers were used: A20, forward 5’-AAACCAATGTTGATTGGAAACTG-3’ and reverse 5’-GTTTGGCTCAATCTGATCC-3’; β-actin, forward 5’-GCTTCTAGCGGACTGTACTGA-3’ and reverse 5’-GCCATGCCAATGTTGCTCTTAT-3’; Bcl-2, forward 5’-GTACCTGAACGGCATCCTG-3’ and reverse 5’-GGGGCCATATAGGGATTGTC-3’; cIAP, forward 5’-TACGGATGAAGGGTCAGGAGT-3’ and reverse 5’-GCCATGCCAATGTTGCTCTTAT-3’; Bim, forward 5’-CCCGGAGTACGATTGC-3’ and reverse 5’-CTCGCCCGTGAATCTTGGC-3’; claf, forward 5’-TACGGATAGGATTTGACTG-3’ and reverse 5’-GCCACCTGCTCCTGTTAGG-3’; GAPDH, forward 5’-GCA TGGCTTCTCCGTGTC-3’ and reverse 5’-GTACCATCATACTTGGAAGTTTC-3’; HPRT, forward 5’-TGAAGGCTACTGTAAATGTACGATCA-3’ and reverse 5’-AGCAAGCTTGGCAACCTATACC-3’; Mcl-1, forward 5’-CAAAGATGGGGTAAACTGCGG-3’ and reverse 5’-CGTTTGCCTTACAAAGA-3’; Noxa, forward 5’-ACTGTTGTTCTGGCGGAGAT-3’ and reverse 5’-TGAGACAACATGGCTCCTTCAA-3’; Puma, forward 5’-ATGGCGGAGACCCTCTAAC-3’ and reverse 5’-GGATGCCCATGGAAGATAG-3’ and XIAP, forward 5’-ATGGCGGAGACCCTCTAAC-3’ and reverse 5’-GAGATGCTGCTGCTGACTG-3’. mRNA expression levels were analyzed using qBase+ software (Biogazelle).

Cell culture and stimulation

For the measurement of IFNγ production and degranulation, 3 × 10⁶ splenocytes were cultured in RPMI 1640 (Gibco) supplemented with 10% FCS (Bodinco) in the presence of GolgiStop (BD) and anti-CD107a (BD). The cells were cultured in the
presence of cytokines (rmIL-12 at 25 ng/ml [Peprotech], rmIL-18 at 5 ng/ml [Peprotech], and rmIL-2 at 20 ng/ml [Protein Service Facility, VIB]) or were cultured on antibody-coated plates (anti-NK1.1 at 10 µg/ml) for 4 h at 37°C. For p-4EBP1 and p-S6 phosphoflow, 4 × 10^6 splenocytes were stimulated for 1 h at 37°C in the presence or absence of rmIL-15 (100 ng/ml; Peprotech), after which cell surface stainings were performed as described, followed by fixation and intracellular stainings. For steady state and stimulation-induced P-p65 phosphoflow, 3 × 10^6 splenocytes were either stained immediately ex vivo or stimulated for 5 or 15 min at 37°C with 1 or 10 ng/ml mouse TNF (mTNF; Tissue Culture Core, VIB). Untreated cells were used as controls at these time points. Culture medium for phosphoflow experiments consisted of RPMI 1640 supplemented with 10% FCS. For TNF hypersensitivity, 2 × 10^6 splenocytes were cultured for 0, 3, 6, 9, and 12 h at 37°C in tissue culture medium (RPMI 1640, 10% FCS, Glutamax [Life Technologies], Gentamicin [Gibco], and β-mercaptoethanol [Tissue Culture Core, VIB]) and supplemented with 0, 1, 2.5, or 5 ng/ml mTNF. 9- and 12-h cultures were supplemented with low-dose rmIL-15 (10 ng/ml; Peprotech). Cell viability was determined by annexin V and live/dead staining, as described. For glucose uptake, 2 × 10^6 splenocytes were incubated for 10 min in RPMI 1640 supplemented with 100 µM of the fluorescent glucose analogue 2-NBDG (Thermo Fisher Scientific) was chemically linked to OP-Puro (Thermo Fisher Scientific) containing 200 µM CuSO₄. Immediately after preparation, cells were incubated with this mixture at room temperature. After 10-min incubation, the reaction was quenched by addition of PBS supplemented with 5% FCS (Bodinco) and 5 mM EDTA (Lonza). Cells were washed twice to remove unbound AF647 PCA.

Measurement of protein synthesis
OP-Puro (Jena Bioscience) was dissolved in DMSO, further diluted in endotoxin-free PBS (50 mg/ml; Gibco), and injected i.p. (50 mg/kg). 1 h after injection, mice were euthanized, and lymphocytes were obtained from the spleens as described above. 5 × 10^6 cells were stained for surface markers and subsequently fixed and permeabilized using the Foxp3 kit (eBioscience). For OP-Puro labeling, Alexa Fluor 647 picolyl azide (AF647-OP; Thermo Fisher Scientific) was chemically linked to OP-Puro through a copper-catalyzed azide-alkyne cycloaddition. In short, 0.5 µM AF647-OP was dissolved in the Click-IT Cell Reaction Buffer (Thermo Fisher Scientific) containing 200 µM CuSO₄. Immediately after preparation, cells were incubated with this mixture at room temperature. After 10-min incubation, the reaction was quenched by addition of PBS supplemented with 5% FCS (Bodinco) and 5 mM EDTA (Lonza). Cells were washed twice to remove unbound AF647-OP.

Statistical analyses
Datasets were analyzed with two-tailed unpaired Student’s t test, Mann–Whitney U test (i.e., nonparametric analogue of Student’s t test), or one-way ANOVA combined with Tukey’s multiple comparisons test. For TNF hypersensitivity, we performed area under the curve (AUC) analysis on normalized data, determined by the ratio of live cells at a given time point (3, 6, 9, or 12 h) to live cells at the start of the experiment (0 h). Ratios were determined per biological replicate per dose. Ad hoc unpaired Student’s t tests were performed on average normalized AUC data per dose per genotype. All statistical tests were performed with Prism (GraphPad Software), and P values express the level of significance (*, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001). Error bars represent the SD.

Online supplemental material
Fig. S1 shows the immunophenotyping of different immune cell populations in spleen, liver, and LP-SI of NK-A20 mice. Fig. S2 shows an overview of apoptotic markers in A20-deficient NK cells and the effect of necroptosis inhibition in an adoptive transfer setting. Fig. S3 describes TNF levels in A20-deficient NK cells and A20- and TNF compound-deficient NK cells in addition to TNF levels in serum and in ex vivo-cultured splenocytes in different A20 mutant mouse lines.

Acknowledgments
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ELISA
TNF secretion was determined by ELISA (eBioscience) following the manufacturer’s guidelines. Samples included serum or supernatant from 24-h–cultured enriched NK cells or splenocytes. See Preparation of single-cell suspensions and Cell culture and stimulations for further details.

Adoptive cell transfers
NK cells were isolated and enriched from the spleens of CD45.1 (WT) or CD45.1.2 (WT) and A20fl/fl × ERT2-Cre+ animals, and a 1:1 mixture of donor cells (total number of cells: 4–5 × 10^7) was given i.v. to congenic recipients. Immediately after cell transfer, recipient mice were treated with tamoxifen (or corn oil as control) for 5 d, and in some cases also with indicated chemical compounds for 6 d consecutively, unless stated otherwise. The ratio of donor NK cells in the blood or spleen of the recipient animals was determined by flow cytometry. For validation of the tamoxifen-inducible ablation of A20, donor NK cells were sorted from the spleens of recipients after 2- or 3-d tamoxifen treatment, and A20 expression was determined by RT-qPCR.

Chemical compounds
Tamoxifen (Sigma-Aldrich) was dissolved in corn oil (20 mg/ml; Sigma-Aldrich) and injected i.p. (2 mg/mouse) for 5 d, unless stated otherwise. Corn oil (Sigma-Aldrich) was used as vehicle control. Nec1s (100 mM in DMSO; kindly donated by N. Takeda) was injected i.v. (6.25 mg/kg) after further dilution in endotoxin-free PBS (Gibco) to reach a final DMSO content of 2.5%. Rapamycin (InvivoGen) was dissolved in DMSO (10 mM), diluted in endotoxin-free PBS (Gibco), and injected i.p. (0.6 mg/kg) at a final DMSO content of 1%. TNFa (2.6 mg/ml; clone XT22; Biocers) and isotype control (rat IgG1; clone GL113; 1.9 mg/ml; Biocers) were diluted in endotoxin-free PBS (Gibco) and injected i.p. (10 mg/kg). Treatment with one of the above compounds started 1 d after the start of the tamoxifen treatment and lasted 6 d.

Statistical analyses
Datasets were analyzed with two-tailed unpaired Student’s t test, Mann–Whitney U test, or one-way ANOVA combined with Tukey’s multiple comparisons test. For TNF hypersensitivity, we performed area under the curve (AUC) analysis on normalized data, determined by the ratio of live cells at a given time point (3, 6, 9, or 12 h) to live cells at the start of the experiment (0 h). Ratios were determined per biological replicate per dose. Ad hoc unpaired Student’s t tests were performed on average normalized AUC data per dose per genotype. All statistical tests were performed with Prism (GraphPad Software), and P values express the level of significance (*, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001). Error bars represent the SD.

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