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Regnase-1 and Roquin regulate a common element in inflammatory mRNAs by spatiotemporally distinct mechanisms

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Summary
Regnase-1 and Roquin are RNA binding proteins essential for degradation of inflammation-related mRNAs and maintenance of immune homeostasis. However, their mechanistic relationship has yet to be clarified. Here we show that although Regnase-1 and Roquin regulate an overlapping set of mRNAs via a common stem-loop structure, they function in distinct subcellular locations: ribosome/endoplasmic reticulum and processing-body/stress granules, respectively. Moreover, Regnase-1 specifically cleaves and degrades translationally active mRNAs and requires the helicase activity of UPF1, similar to the decay mechanisms of nonsense mRNAs. In contrast, Roquin controls translationally inactive mRNAs, independent of UPF1. Defects in both Regnase-1 and Roquin lead to large increases in their target mRNAs, although Regnase-1 tends to control the early phase of inflammation when mRNAs are more actively translated. Our findings reveal that differential regulation of mRNAs by Regnase-1 and Roquin depends on their translation status and enables elaborate control of inflammation.
Introduction

Inflammation is mediated by proinflammatory cytokines such as tumor necrosis factor (TNF) and Interleukin 6 (IL-6). Expression of cytokines is rapidly induced in response to infection by pathogens via Toll-like receptors (TLRs) in innate immune cells (Chovatiya and Medzhitov, 2014; Moresco et al., 2011; Takeuchi and Akira, 2010). Whereas cytokine mRNA levels are controlled at both transcriptional and post-transcriptional levels, post-transcriptional dampening of protein expression in particular can resolve inflammation and prevent unintended tissue damage (Anderson, 2010; Kafasla et al., 2014). Eukaryotic mRNAs are in dynamic equilibrium between different subcellular locations: actively translated mRNAs can be found in polysomes, mRNAs stalled in translation initiation can accumulate in stress granules (SGs), and mRNAs targeted for degradation or translation repression can accumulate in processing bodies (PBs). Further, mRNA-protein (mRNP) complexes dynamically move between polysomes, SGs and PBs (Anderson and Kedersha, 2009; Balagopal and Parker, 2009).

Many immune-related mRNAs have short half-lives because of conserved cis-elements including AU-rich elements (AREs) and stem-loop (SL) structures in their 3’ untranslated regions (UTRs) (Anderson, 2010; Kafasla et al., 2014). ARE binding proteins are involved in controlling the stability of such mRNAs. In addition, SLs present in mRNAs including ICOS, OX40 and TNF are destabilized by Roquin-1 and -2 (Leppek et al., 2013). These proteins harbor an RNA binding ROQ domain whose loss of function mutation (M199R) in mice (San) leads to the development of autoimmune disease characterized by an increase in follicular helper T cells (Linterman et al., 2009; Vinuesa et al., 2005; Yu et al., 2007). ARE- and Roquin-mediated mRNA decay takes place in PBs or SGs where a CCR4-CAF1-NOT deadenylase complex is recruited (Anderson, 2010; Athanasopoulos et al., 2010).

In contrast, another set of mRNA decay pathways target mRNAs undergoing translation. In the nonsense mediated decay (NMD), premature translation termination
at more than 20-25 bases 5′ of the exon junction complex (EJC) promotes the recruitment of an SMG1-UPF1-eRF1-eRF3 (SURF) complex (Kervestin and Jacobson, 2012; Popp and Maquat, 2013; Schweingruber et al., 2013). The SURF complex then interacts with the UPF2-UPF3-EJC complex, leading to recruitment of the endonucleases SMG6, SMG5 and SMG7, which mediate deadenylation for further degradation. In addition, active translation and UPF1 are required for some normal mRNA decay pathways, including staufen1-mediated mRNA decay (SMD) and replication-dependent histone mRNA decay (Kaygun and Marzluff, 2005; Kim et al., 2005).

We identified Regnase-1 (Reg1; also known as Zc3h12a and Mcpip1) as an RNase critical for preventing a severe autoimmune inflammatory disease in mice by destabilizing inflammation-related mRNAs (Iwasaki et al., 2011; Matsushita et al., 2009). Reg1 harbors a PIN-like RNase domain, and controls a set of genes including \textit{Il6} and \textit{Reg1} itself, but not \textit{Nfkbia}, in macrophages following stimulation with TLR ligands such as lipopolysaccharide (LPS) and IL-1β, which activate shared signaling pathways. Reg1 is also essential for suppressing aberrant activation of T cells by targeting genes such as \textit{Icos}, \textit{c-Rel} and \textit{Ox40} for degradation (Uehata et al., 2013).

In this study, we aim to uncover how Reg1 recognizes and degrades its target mRNAs, and show that Reg1 interacts with a set of SL-containing mRNAs that overlaps with those targeted by Roquin. In contrast to Roquin, Reg1 co-localizes with ribosomes, but not with PBs and SGs. Reg1 destabilizes translationally active mRNAs, and requires UPF1. The apparent cooperation between Reg1 and Roquin is confirmed by the upregulation of their mutual target mRNAs under mutation of both Reg1 and Roquin, although Reg1 and Roquin tend to control the early and late phases of inflammation, respectively. Thus, a common set of SL-containing mRNAs are recognized by distinct proteins in different stages of mRNA metabolism for fine tuning of immune responses.
Results

SL structure required for Reg1-RNA association.

We previously showed that a potential SL sequence in the *Il6* 3′UTR (84-102) is recognized by Reg1 for degradation (Matsushita et al., 2009). To determine the secondary structure of the *Il6* 3′UTR target sequence, we performed selective 2′-hydroxyl acylation analyzed by primer extension (SHAPE) analysis. The SHAPE results were consistent with the computational prediction of a simple SL structure with bulge (Figure 1A and 1B). We then examined if the SL is critical for the control of *Il6* expression by Reg1 by using an antisense morpholino that can interfere with SL formation in *Il6* 3′UTR (Il6-SL-MO) (Figure 1C). A luciferase assay expressing the reporter construct with the *Il6* 3′UTR, the morpholinos, and Reg1 revealed that the suppression of the luciferase activity observed in control (Ctrl) morpholino transfected cells was cancelled by the Il6-SL-MO (Figure 1D). Following introduction of the Il6-SL-MO to mouse bone marrow macrophages (BMMs), LPS-induced expression of mRNA and protein for IL-6, but not TNF, were significantly elevated compared with control cells (Figure 1E). We further supplemented *Il6*−/− mice with wild-type (WT) BMM treated with the Il6-SL-MO. LPS-induced production of IL-6, but not TNF, in the sera was elevated in mice supplemented with Il6-SL-MO-treated BMM, indicating that the SL in the *Il6* 3′ UTR is important for the regulation of IL-6 production in vivo (Figure 1F).

To investigate if the conserved SL is required for processing by Reg1, which harbors endonuclease activity as shown by the digestion of circular RNA in vitro (Figure S1A and S1B), we synthesized *Il6* 3′UTR RNAs with and without the SL (Figure S1C). Reg1 degraded RNAs harboring the SL, but not without the SL (Figure S1C), suggesting that the SL is required for RNase-mediated digestion or RNA binding. To distinguish between these two possibilities, we artificially tethered Reg1 to *Il6* coding and luciferase mRNAs using the λN-BoxB system (Figure 1G). Interestingly, a
λN-Reg1, but not WT Reg1, destabilized the Il6 coding sequence in the presence of 5XBoxB in the Tet-off system (Figure 1H and 1I). This was further confirmed by a luciferase reporter assay expressing the luciferase construct with 5XBoxB in the 3’ UTR and λN-Reg1 (Figure S1D), indicating that the SL is required for the recruitment of target mRNAs by Reg1, but is dispensable for their degradation.

To check the possibility of Reg1-mediated control of miRNA generation (Suzuki et al., 2011), we prepared small RNAs from Reg1–/– mouse embryonic fibroblasts (MEFs). QPCR analysis revealed that the levels of miRNAs previously reported to be regulated by Reg1 as well as a set of LPS-inducible miRNAs including miR-155 and miR-146a did not increase in the absence of Reg1 (Figure S1E), indicating that Reg1 is dispensable for the control of these miRNAs.

Identification of mRNAs associating with Reg1.

To comprehensively identify mRNAs associated with Reg1, we performed RNA-immunoprecipitation (IP) sequencing (RIP-Seq) analysis. We expressed Flag-Reg1 lacking RNase activity (D141N), which was predicted to retain RNA binding activity (Figure S1F), in HeLa cells. We found that Il6 and Reg1 (ZC3H12A) were highly enriched by IP with Reg1 compared with control IP, both in unstimulated and IL-1β-stimulated cells (Figure 1J and Table S1), indicating that IP with Reg1 properly co-precipitated its target mRNAs. A total of 68 mRNAs were significantly enriched either in unstimulated or IL-1β-stimulated cells (p values < 10⁻³) compared with control cells (Table S1). Gene ontology annotations enriched for the 68 Reg1-interacting mRNAs were often involved in inflammation (Table S2).

We then investigated if the Reg1-binding mRNAs were targeted by Reg1 for degradation. We constructed luciferase reporter genes with 3'UTRs from a set of enriched genes including Nfkbi, Nfkbid, Ptgs2, Mafk, Id1, Cxcl1, Cxcl2 and Cxcl3. Overexpression of Reg1 suppressed the luciferase activity in an RNase-activity
dependent manner (Figure 1K). Consistent with a previous report (Iwasaki et al., 2011), overexpression of Reg1 suppressed Reg1 3’ UTR containing nucleotides 1-210, but not in the case where nucleotides 1-200, which contains the SL, were absent. Collectively, Reg1-bound mRNAs obtained by RIP-Seq are indeed targeted by Reg1 for degradation.

The Reg1 target consensus motif overlaps with that of Roquin.

Roquin-1 was also reported to associate with SL containing mRNAs such as Tnf, Icos, Nfkbid and Nfkbiz in RAW264.7 cells (Leppek et al., 2013). Indeed, overexpression of Roquin suppressed the luciferase activity in the presence of the 3’UTR of Nfkbiz, Nfkbid and Ptgs2 (Figure S1G), suggesting an overlap between Reg1 and Roquin target mRNAs.

We therefore next examined Roquin-interacting mRNAs globally in HeLa cells by RIP-Seq analysis. A total of 52 high-confidence mRNAs were enriched by IP with Flag-Roquin either in unstimulated or IL-1β-stimulated cells (p values < 0.01) (Table S3). Gene Set Enrichment Analysis (GSEA) showed that 68 Reg1 targets (p ≤ 1.0 X 10^{-3}) were significantly biased toward high enrichment scores (ES) in the Roquin RIP-seq data (P < 1 X 10^{-4}; Figure 2A). We found a similar significant enrichment of Reg1 targets among mRNAs enriched in the Roquin RIP-seq sample when using the Kolmogorov-Smirnov test (P < 2.2 X 10^{-16}). Reciprocally, the set of 52 high-confidence Roquin targets (p ≤ 0.01) showed a significant overlap with mRNAs enriched in the Reg1 RIP-seq data (P < 1 X 10^{-4}; Figure 2B). These 52 Roquin targets shared 10 mRNAs with the 68 Reg1 targets (P = 2.69 X 10^{-15}). These results demonstrate that the Reg1 and Roquin target mRNAs overlap significantly.

Among newly identified Reg1-associated mRNAs, we focused on two mRNAs, Nfkbiz and Ptgs2. By testing luciferase constructs with truncated Nfkbiz 3’UTRs (Figure S2A), we found that Nfkbiz (100-150), which is evolutionally conserved and predicted to contain 2 SLs (Figure S2B and S2C), is required for Reg1-mediated suppression.
Addition of either one of these SL to the β-globin 3'UTR conferred responsiveness to Reg1 (Figure S2D). Similarly, we found that *Ptgs2* (1000-1300) was required for the regulation by Reg1 (Figure S2E). Mouse *Ptgs2* (1211-1228) was predicted and shown to form a conserved SL (Figure S2F-S2H), and this sequence alone was sufficient for Reg1 to suppress luciferase activity (Figure S2I). In addition, we found that a luciferase construct with the constitutive decay element (CDE) present in *Tnf* mRNA, which was identified as a Roquin target SL (Leppek et al., 2013), and SL alone (TNF-CDE37) was suppressed by Reg1 (Figure 2C and S2J).

Next, we investigated target structures present in Reg1 binding mRNAs globally by high-throughput sequencing of RNA isolated by crosslinking immunoprecipitation (HITS-CLIP) in HEK293 cells (chosen because of their wide usage in CLIP analysis) with stable, inducible expression of Flag-tagged WT Reg1 (Figure 2D). UV-crosslinked RNAs co-precipitated with Flag-Reg1 were recovered and sequenced. By analyzing sequence data of two biological replicate libraries, we identified 9107 and 10448 putative Reg1 binding groups in two biological duplicate libraries, respectively (Table S4). The two libraries intersected on a group level with at least one overlapping nucleotide; we thereby obtained 1143 Reg1 binding sites. As expected, many sites were located in 3'UTRs, although some fractions were mapped to coding regions and introns (Figure 2E). Although immune-related mRNAs were not frequently observed in HEK293 cells because of their poor expression, we identified Reg1 target sites in genes such as TM2D3, which was identified by RIP-seq analysis. The obtained sequences from them were predicted to form SLs (Figure 2F), and were suppressed by Reg1 (Figure 2G). When the Reg1 binding sequences were globally folded using RNAfold, SL structures with 3-7 nucleotide stems and 3 nucleotide loops were significantly enriched compared with 1000 individual permutations in both of the replicates and their intersection (p < 0.001) (Figure 2H). Next, we investigated the sequence motifs enriched in the Reg1 binding SLs harboring a hairpin of 3 nucleotides.
with varying stem lengths (3, 5 or 7 nucleotides). Interestingly, the UAU sequence was significantly enriched in the hairpin sequences (Figure 2I).

Next, we examined the requirement of the SL structure and the specific hairpin loop sequence for Reg1-mediated target mRNA suppression. Using two validated SLs, TNF-CDE and another artificial SL with an UAU loop (Figure S3A-S3C), we first analyzed the importance of SL structures. Disruption of the SL structures abrogated Reg1-mediated suppression, and restoration of the SL structure by mutating both sides of the stem resulted in rescue of Reg1-mediated suppression (Figure S3D and S3E). We then generated a set of luciferase constructs harboring SLs with different loop sequences. In addition to SLs with UAU, UGU loop sequences were significantly inhibited by Reg1 overexpression (Figure 2J), indicating that SL sequences with either A or G at the second residue in the hairpin loop can be suppressed by Reg1. In contrast, ACA, AAA or UCU SLs were not suppressed by Reg1 expression, suggesting that a pyrimidine-purine-pyrimidine (Py-Pu-Py) sequence should be present in the loop.

Furthermore, IP with Reg1 co-precipitated reporter mRNA with a SL structure, while reporters whose SL was disrupted no longer co-precipitated with Reg1 (Figure S3F and S3G), indicating that the SL structure is required for Reg1 binding in cells. Additionally, alteration in the loop sequence abrogated its binding with Reg1 (Figure S3H).

Reciprocally, Roquin overexpression suppressed the luciferase activity and mRNA with β-globin 3’UTR followed by Reg1 target SL sequences (Figure 2K). Taken together, these results demonstrate that Reg1 and Roquin can recognize overlapping target mRNAs via the same SLs present in their 3’UTRs.

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**Reg1 localizes to the endoplasmic reticulum (ER), but not PBs or SGs.**

The fact that Reg1 and Roquin recognize the same SL prompted us to investigate their relationship further. We first examined the involvement of deadenylation and decapping
in Reg1-mediated mRNA decay by overexpressing dominant negative mutants of Caf1 (Caf1-AA), and Dcp2 (Dcp2-AA). In contrast to Roquin-1 and -2, neither Caf1-AA nor Dcp2-AA affected Reg1-mediated degradation of *Il6* mRNA (Figure S4A).

We next investigated if Reg1 and Roquin function in the same subcellular location. Immunofluorescence staining revealed that Roquin-1 localizes in PBs in NIH3T3 and HeLa cells and in SGs upon arsenite treatment (Figure S4B and S4C). In contrast, NIH3T3 cells expressing Flag-Reg1 did not localize to PBs, or SGs upon arsenite treatment (Figure 3A and Figure S4D). In contrast, Reg1 co-localized with Calnexin, a type I membrane protein of the ER, but not with disulfide isomerase (PDI), a protein found within the ER (Figure 3B and S4E), indicating that Reg1 localizes on the surface of the ER. Because the surface of the rough ER (RER) is studded with ribosomes, we next demonstrated that Reg1 also co-localizes to the ribosome (Figure 3B and S4E). Moreover, immunoelectron microscopy analysis revealed that Reg1 localized to the cytoplasm and the surface of the ER (Figure 3C). Furthermore, subcellular fractionation of NIH3T3 cells revealed that endogenous Reg1 was present in the cytosol and RER, but not smooth ER, which does not contain ribosomes (Figure 3D).

To further investigate whether Reg1 localizes to translationally active polysomes, HeLa cell lysates were subjected to polysome fractionations (Figure 3E). Immunoblot analysis revealed that endogenous Reg1 localized in polysomal fractions in addition to non-polysome (non-ribosome and 40S-80S) fractions (Figure 3E), though Roquin-1 and -2 were localized in the non-polysome fractions.

**Reg1, but not Roquin, regulates target mRNAs in the polysomal fraction.**

Next we extracted RNAs from ribosomal fractions in HeLa cells following Reg1 and Roquin-1/-2 siRNA treatment with or without IL-1β stimulation, and examined the levels of mRNA expression in each fraction (Figure 3F and 3G). Reg1 and Roquin-1/-2
were efficiently suppressed at the mRNA and protein levels (Figure 3G and data not shown). Knockdown of Reg1 resulted in a large increase in IL6 and PTGS2 mRNA expression in the polysome fractions (fractions 7-12) compared with control cells, whereas the mRNA in non-polysome fractions (fractions 3-6) were comparable (Figure 3H). However, TNF and NFKBIA expression was unchanged in Reg1 knockdown. In contrast, TNF and PTGS2 increased in Roquin-1/-2 knockdown cells. Interestingly, the increase was observed in non-polysome fractions, but not in polysome fractions (Figure 3H).

We further detected PTGS2 in HeLa cells by using in situ hybridization following IL-1β stimulation (Figure S4F). Knockdown of Reg1 and Roquin-1/2 resulted in an increase in PTGS2 expression (Figure S4F and S4G). Whereas Reg1 knockdown decreased PTGS2 localized in the PB, Roquin-1/-2 knockdown increased PTGS2 localized in PBs (Figure S4F and S4H). Taken together, these results demonstrate that Reg1 is critical for controlling the expression of translationally active mRNAs, whereas Roquin-1/2 regulate translationally inactive mRNAs in PBs/SGs.

**Reg1 destabilizes translationally active mRNA via stop codons.**

We next treated Reg1−/− macrophages with protein translation inhibitors, anisomycin and cycloheximide (CHX), and examined the kinetics of Il6 and Ptgs2 mRNA degradation after stimulation with LPS. Whereas Il6 and Ptgs2 mRNAs were stabilized in Reg1−/− macrophages compared with WT, the difference was lost following treatment with anisomycin or CHX (Figure 4A). Reciprocally, destabilization of Il6 mRNA induced by Reg1 overexpression was no longer observed in response to treatment with these protein synthesis inhibitors (Figure 4B and 4C). The translation inhibitors did not alter Reg1 cellular localization or its protein expression levels (Figure S5A and S5B).

We next examined the effect of Reg1 on translationally inactive mRNA. We inserted a highly stable SL (SSL), which was shown to inhibit translation (Doma and
Parker, 2006), immediately upstream of the start codon of luciferase or \textit{Il6} (Figure 4D). Insertion of the SSL to the luciferase reporter resulted in severe decrease in luciferase activity, but not in its mRNA, suggesting that the SSL suppressed protein synthesis (Figure 4E). Overexpression of Reg1 no longer destabilized \textit{Il6} with the SSL in the 5′UTR (Figure 4F and 4G), further confirming that active translation is essential for Reg1-mediated mRNA decay.

However, Reg1-mediated mRNA destabilization was not affected by the presence of IRES at the 5′UTR (Figure S5C and S5D), suggesting that translation initiation is dispensable for the function of Reg1. In contrast, Reg1 failed to destabilize an \textit{Il6} whose stop codons were mutated throughout the 3′UTR (\textit{Il6} mRNA-ΔSTOP) and luciferase \textit{Il6} 3′UTR without a stop codon (Figure 4F-4H). Requirement of stop codons in the Reg1-mediated mRNA decay prompted us to hypothesize that the distance between the stop codon and the \textit{Il6} mRNA SL is important for Reg1–mediated destabilization. To examine this, we prepared \textit{Il6} constructs harboring different distances between the stop codon and the SL (Figure 4I). Reg1 destabilized \textit{Il6} mRNA with its original 3′UTR with (1-125), but not without (1-81), a SL (Figure 4J). Although the construct harboring the SL 23 bases downstream of the stop codon (64-125) was still suppressed by Reg1, those with the SL immediately downstream of the stop codon (82-125) were not (Figure 4J). These data suggest that the stop codon and SL structure require a minimal distance (about >20 nt) to be suppressed by Reg1.

\textbf{Reg1 associates with ribosome proteins and UPF1.}

Next we explored Reg1-associated proteins with a global proteomics approach using iTRAQ (Figure 5A). Levels of many ribosomal proteins in Reg1 immunoprecipitates from HeLa and RAW cells expressing Flag-Reg1 increased compared to control cells (Figure 5B and Table S5). Consistently, recombinant Reg1 precipitated by ultracentrifugation when incubated with highly-purified ribosome \textit{in vitro} (Figure 5C),
suggesting that Reg1 is physically associated with the ribosome.

In addition to ribosomal proteins, we found that Flag-Reg1 co-precipitated UPF1 irrespective of Benzonase treatment (Figure 5B and 5D), suggesting that Reg1 and UPF1 exhibit direct protein-protein interaction. Whereas Reg1 lacking the N-terminal helical domain (90-596) (Figure 5E) was able to bind UPF1 (Figure 5F), Reg1 including the RNase domain to the C-terminal end (130-596) or only the C-terminal region (331-596) failed to co-precipitate UPF1 (Figure 5F). In contrast, Reg1 lacking the linker region between the N-terminal helical domain and the RNase domain (Δ90-130) showed severely impaired interaction with UPF1 (Figure 5F). Furthermore, overexpression of Reg1 (Δ90-130) lost the competency to suppress Il6 and Ptgs2 3′UTR (Figure 5G), suggesting that the interaction between Reg1 and UPF1 is essential for Reg1 function.

**UPF1 is critical for Reg1-mediated mRNA decay via its helicase activity.**

Next we examined if UPF1 is required for Reg1-mediated decay. When we knocked down UPF1 in HEK293 cells, overexpressed Reg1 failed to degrade Il6 (Figure 6A and 6B). Furthermore, Reg1 overexpression no longer suppressed the luciferase reporter constructs harboring 3′UTRs for IL6, PTGS2, ICOS and TNF-CDE37 in HeLa and RAW264.7 cells (Figure 6C and Figure S6A and S6B). In contrast, knockdown of UPF2 and UPF3x did not affect Reg1-mediated suppression (Figure 6C and Figure S6B). Further, the response to Roquin overexpression was not inhibited by the loss of UPF1 (Figure 6D and Figure S6C), indicating that Reg1 and Roquin degrade target mRNA in UPF1-dependent and –independent manners, respectively. Interestingly, knockdown of UPF1 or Reg1 in HeLa and RAW264.1 cells resulted in increased expression of IL6 and PTGS2, but not NFKBIA, in response to IL-1β or LPS stimulation, respectively (Figure 6E and S6D). Knockdown of UPF1, but not Reg1, resulted in an increase in NMD-target genes such as SMG5 and GAS5 (Figure 6E and S6D).
Whereas overexpressed Reg1 co-precipitated *Il6*, UPF1 knockdown increased the amounts of *Il6* co-precipitated with Reg1 (Figure 6F). Knockdown of UPF1 abrogated Reg1-mediated inhibition, even when we tethered Reg1 to the target mRNA (Figure S6E and Figure 6G), indicating that UPF1 is required for Reg1 to cleave its target mRNAs after binding. Furthermore, anisomycin treatment abrogated the λN-Reg1-mediated suppression of *Il6*-5XBoxB expression (Figure 6H) without inhibiting the Reg1-*Il6* interaction (Figure S6F), indicating that translation is required for Reg1 to degrade associated mRNAs. The interaction between Reg1 and UPF1 was inhibited by the treatment with anisomycin (Figure 6I), suggesting that the association between Reg1 and UPF1 is mediated by translation.

We next examined the role of UPF1 helicase activity in controlling Reg1. We depleted endogenous UPF1 with siRNA and reconstituted it with exogenous siRNA-resistant WT and helicase inactive (D648A/E649A; DEAA) (Franks et al., 2010) mutant UPF1 in HeLa cells (Figure 6J). WT, but not the DEAA, rescued Reg1-mediated suppression of *Il6* 3'UTR (Figure 6K). Furthermore, IL-1β-mediated expression of *IL6* and *PTGS2* was suppressed by the UPF1 reconstitution in a manner dependent on the helicase activity (Figure 6L). Nevertheless, the UPF1 DEAA was capable of interacting with Reg1 (Figure S6G), indicating that the UPF1 helicase activity is required for Reg1 at the degradation step of its target mRNAs. Collectively, these results demonstrate that UPF1 interacts with Reg1 upon translation, and is essential for Reg1-mediated degradation of target mRNAs after recognition by acting as an RNA helicase.

**Cooperative and distinct roles of Reg1 and Roquin in inflammatory gene expression.**

Since Reg1 and Roquin suppress common target mRNAs via distinct molecular mechanisms, we investigated their relationship by expressing mutant Reg1 and Roquin in the same cells. First, we observed that the Reg1 D141N localized in SGs in addition...
to ER following arsenite treatment (Figure S7A). Interestingly, the Reg1 D141N mutant cancelled suppression of luciferase reporter constructs harboring 3′UTRs for *Il6* (86-102) and *Ptgs2* 3′UTR by Roquin overexpression (Figure S7B). In contrast, Roquin-NT, whose localization is consistent with WT Roquin (data not shown), failed to alter Reg1-mediated suppression (Figure S7B). These results suggest that Reg1 and Roquin affect the same SL, and that the Reg1 D141N mutant stays on the 3′UTR and thereby inhibits the action of Roquin.

We further examined the roles of Reg1 and Roquin by generating BMMs harboring *Reg1*−/− and *Roquin-1*San/San alleles. We found that the expression of *Il6* and *Tnf* upon LPS stimulation was augmented in *Reg1*−/− and *Roquin-1*San/San BMM, respectively (Figure 7A). Further, *Ptgs2* mRNA increased in *Reg1*−/− and *Roquin-1*San/San BMM. Since *Il6*, *Tnf* and *Ptgs2* mRNAs present in polysomal fractions rapidly increased in response to IL-1β stimulation in HeLa cells and then decreased at later time points (Figure 3H and S7C), we hypothesized that the contribution of translation-dependent decay by Reg1 changes in the time-course of BMM activation. Indeed, the difference in *Il6* and *Ptgs2* expression between WT and *Reg1*−/− BMM was more obvious 1 or 2 h after LPS stimulation than 8 h post stimulation (Figure 7A). In contrast, the difference in *Tnf* and *Ptgs2* expression in WT and *Roquin-1*San/San BMM was greater 4 and 8 h after LPS than at earlier time points. These data demonstrate that Reg1 and Roquin play more important roles earlier and later, respectively, in response to TLR stimulation. Consistent with the *Ptgs2* mRNA expression results, Ptgs2 protein was expressed more abundantly in *Reg1*−/− BMM 2 h after LPS stimulation compared with WT (Figure 7B). In contrast, the expression of Ptgs2 protein in *Roquin-1*San/San BMM increased greatly 8 h after stimulation. Furthermore, the Ptgs2 protein/mRNA ratio was higher in *Reg1*−/− BMM compared with WT and *Roquin-1*San/San cells (Figure 7C). These results are consistent with the increased expression of *Ptgs2* mRNAs in polysomal fractions in HeLa cells with defective expression of Reg1 (Figure 3H).
Although Reg1<sup>−/−</sup>; Roquin-1<sup>San/San</sup> double mutant mice were mostly embryonic lethal (data not shown), we successfully established MEFs with the Reg1<sup>−/−</sup>; Roquin<sup>San/San</sup> genotype. In response to LPS and TNF stimulation in WT MEFs, the expression of genes including Il6, Tnf and Ptgs2 was upregulated (Figure 7D and S7D). MEFs lacking Reg1 showed increased expression of Il6 and Ptgs2 compared with WT, particularly in the early phase of the responses (Figure 7D). Similarly, the Roquin San mutation resulted in an increase in the expression of Tnf, Il6 and Ptgs2 at later time points after LPS stimulation (Figure 7D). Interestingly, Reg1<sup>−/−</sup>; Roquin<sup>San/San</sup> MEFs showed high increases in the expression of Il6, Tnf and Ptgs2 mRNAs both in early and late phases, suggesting that Reg1 and Roquin in part redundantly regulate their target mRNAs, although Nfkbia expression was not altered (Figure 7D and S7D).

Finally, we examined the genome-wide changes in gene expression in WT, Reg1<sup>−/−</sup>, Roquin-1<sup>San/San</sup> and Reg1<sup>−/−</sup>; Roquin-1<sup>San/San</sup> MEFs with or without LPS stimulation by transcriptome analysis (Table S6). A set of genes was greatly increased in Reg1<sup>−/−</sup>; Roquin-1<sup>San/San</sup> double mutant MEFs compared with WT (Figure S7E). GSEA revealed that the mouse counterparts of Reg1- and Roquin-binding mRNAs identified by RIP-sequencing analysis were significantly enriched in genes more abundantly expressed in Reg1<sup>−/−</sup>; Roquin-1<sup>San/San</sup>, Reg1<sup>−/−</sup> and Roquin-1<sup>San/San</sup> MEFs (Figure 7E, 7F and Figure S7F). These results indicate that the Reg1- and Roquin-binding mRNAs are commonly regulated by these RNA binding proteins.
Discussion

In this study, we found that Reg1 and Roquin target overlapping mRNAs and SLs for degradation. Nevertheless, they degrade the target mRNAs using different molecular mechanisms and in different cellular locations. Whereas Roquin localizes in PBs/SGs and degrades mRNAs via deadenylation like other mRNA decay pathways, Reg1 associates with ribosomes, and degrades mRNAs in a translation-dependent manner (Figure 7G).

The UAU loop sequence was enriched in Reg1-binding sites, and the loop sequence was important for Reg1-mediated mRNA decay. So far, all validated Reg1 target SLs contain a Py-Pu-Py loop sequence, and mutation of this loop sequence results in Reg1 unresponsiveness, a rule shared by Roquin target mRNAs. We did not find a specific rule in the stem-sequence of Reg1-target mRNAs, although Roquin target stem sequences were reported to harbor the UUG sequence in their upper stem. One possibility is that Reg1 can recognize a broader range of SL structures than Roquin. However, crystal structures studies revealed that Roquin ROQ domain interacts with the TNF-CDE stem and a triloop with Py-Pu-Py sequences, and that Roquin uses mainly non-sequence-specific contacts with the RNA (Schlundt et al., 2014; Schuetz et al., 2014; Tan et al., 2014). Indeed, we found that overexpression of Roquin could affect 3′ UTRs harboring a stem without the UUC sequence. Therefore, the requirement for particular stem sequences may not be so strict, and SL structures harboring a Py-Pu-Py loop sequence might be more broadly recognized by both Reg1 and Roquin.

Reg1 and Roquin negatively regulate activation of T cells in addition to the control of inflammatory gene expression in innate immune cells (Pratama et al., 2013; Uehata et al., 2013; Vogel et al., 2013). Icos is a common target of Reg1 and Roquin proteins, and we found that a SL structure present in its 3′UTR is recognized by both Reg1 and Roquin. A recent report showed that Nfkbiz and Nfkbid are suppressed by Reg1 and Roquin in T cells, controlling Th17 differentiation, although their relationship
was not studied (Jeltsch et al., 2014). We focused on the mechanisms in innate immune and non-immune cells in this study; however, it is likely that Reg1 and Roquin similarly regulate mRNAs related to T cell activation in a manner dependent on the state of translation. Since lethality upon mutation of both Reg1 and Roquin prohibited analyzing their roles in adult T cells, it would be interesting to analyze their functional roles in T cells by using conditional alleles in the future.

We found that Reg1-mediated mRNA decay requires translation and translation termination approximately 20 bases or more upstream of the SL. Similar to NMD, the stop codon might be required for preventing the disruption of the SL by the ribosome. In addition, the requirement of UPF1 demonstrates that the Reg1-mediated decay mechanism is highly similar to that of the quality control pathway, NMD. In the execution of NMD, endonucleolytic cleavage by SMG6 as well as exonucleolytic cleavage induced by the complex of SMG5-SMG7 contributes to the degradation of aberrant mRNAs (Eberle et al., 2009; Loh et al., 2013). SMG6-cleaved mRNA product can be further degraded in a manner dependent on the exosome and XRN1 exonucleases. Since Reg1 serves as an endonuclease like SMG6, a similar complete mRNA degradation system might operate upon cleavage by Reg1.

An open question is why Reg1-mediated mRNA decay requires active translation. Given that translation-inactive mRNAs are not affected by the absence of Reg1, this mechanism may be beneficial for maintaining a certain reserve of inflammation-related mRNAs. If Reg1 were to arbitrarily degrade its target mRNAs irrespective of their translation status, immune-related mRNA levels might become very low and additional energy would be required for de novo transcription. Alternatively, this mechanism may lead to the production of one protein per mRNA to keep control of short half-lives of Reg1-target mRNAs. In contrast, Roquin-mediated decay is involved in the constitutive decay of mRNAs and can be important for the regulation of the amounts of stored immune-related mRNAs. The presence of different regulatory
proteins makes posttranslational control of inflammation more elaborate than previously realized.

Although UPF1 is essential for NMD, the molecular mechanisms how UPF1 is involved in NMD are still under debate. In one model, UPF1 is recruited to the mRNP complex upon translation termination forming a SURF complex (Kurosaki and Maquat, 2013), and the helicase activity of UPF1 is proposed to be required for the disassembly of the mRNP complex (Franks et al., 2010). However, this model was challenged in reports demonstrating that UPF1 is associated with mRNAs even before translation (Hogg and Goff, 2010; Hurt et al., 2013; Zund et al., 2013). The helicase activity of UPF1 is essential for Reg1-mediated cleavage of target mRNAs, although this activity is dispensable for Reg1 to interact with its target mRNAs. Therefore, the Reg1-UPF1 interaction is possibly induced in the course of translation irrespective of UPF1 helicase activity, which may potentiate the RNase activity of Reg1 by remodeling an mRNP complex (Figure 7G).

Reg1-mediated decay is dependent on UPF1, but not on UPF2 and UPF3x. Whereas all three UPF members are required for NMD, UPF2 and UPF3x are also dispensable for SMD and replication-dependent histone mRNA decay (Kaygun and Marzluff, 2005; Kim et al., 2005). Nevertheless, Staufen 1 and 2 are sensors for recognizing dsRNA sequences (Park et al., 2013), but do not have RNase activity. Staufen1 was dispensable for the Reg1-mediated decay (data not shown), and Reg1-mediated mRNA decay is highly unique in that Reg1 serves as both a sensor for the SL and a degradation enzyme simultaneously.

Although loss of function of both Reg1 and Roquin-1/-2 resulted in a large increase in the expression of Tnf, Il6 and Ptgs2, dysfunction of Reg1 or Roquin alone seems to differentially regulate Il6 and Tnf, respectively. Further, both Reg1 and Roquin-1 suppressed Tnf and Il6 mRNA 3′ UTRs upon overexpression; thus, it is currently difficult to explain why Tnf expression did not increase under Reg1 deficiency.
Since Reg1 is critical for the regulation of \( \text{Tnf} \) in the absence of Roquin, Roquin may bind with the \( \text{Tnf} \) SL more strongly than Reg1 and thus reduce the accessibility of Reg1 to \( \text{Tnf} \) mRNA. Furthermore, it would not be surprising if the translation status dynamically changes in the course of the life of an mRNA. Further studies will uncover the mechanisms of dynamic cellular regulation of mRNA translation status in relation to mRNA decay.

Collectively, this study clearly demonstrates that the posttranscriptional regulation of inflammation is controlled by Reg1 and Roquin in spatiotemporally distinct manners. Given that excess and prolonged production of cytokines leads to the onset of inflammatory diseases, prolonged stability of inflammatory cytokine mRNAs can be considered aberrant, and thus targeted by the quality control system. Differential regulation of Reg1- and Roquin-mediated mRNA degradation are thus necessary for the elaborate control of inflammation.
Experimental Procedures

RIP-sequencing
HeLa expressing Flag-Reg1-D141N, Roquin-1 or an empty vector were stimulated with human IL-1β (10 ng/mL) for 2 h. RNA-protein complexes were immunoprecipitated with anti-Flag antibody (F3165; Sigma) bound to protein G magnetic beads and RNAs were extracted from the beads. RNA library was prepared using Small RNA Sample Prep kit v1.0 (Illumina Inc.) and sequenced on a HiSeq 2000 system (Illumina) also according to the manufacturer’s instructions. The resulting set of trimmed reads were then mapped against the human genome (hg19; NCBI).

mRNA decay assay
For mRNA decay experiments with Tet-off system, doxycycline (1 μg/ml) (Sigma) was added to the medium for the indicated time intervals before harvesting Tet-off 293 cells. Total RNA was isolated using Trizol (Invitrogen). mRNA levels were determined either by northern blot or by RT-qPCR analysis. For determining RNA decay using the Click-iT® Nascent RNA Capture Kit (Invitrogen), peritoneal macrophages were labelled with EU (0.5 mM) and stimulated with LPS (100 ng/mL) for 2 hr and the medium was replaced with growth medium without EU for the indicated time intervals before harvesting EU-labeled peritoneal macrophages. EU-labeled RNAs (EU-RNAs) were biotinylated and the biotinylated RNAs were purified with streptavidin magnetic beads and analyzed with RT-qPCR.

Luciferase assay
HeLa cells were transfected with luciferase reporter plasmid pGL3 containing the 3′ UTR of indicated genes, together with expression plasmid for Regnase-1 or empty (control) plasmid. After 24 h of cultivation, cells were lysed and luciferase activity in lysates was determined with the Dual-Luciferase Reporter Assay system (Promega).
The gene encoding Renilla luciferase was transfected simultaneously as an internal control.

**Statistical analysis**

Data are presented as means ± standard deviation. Statistical significance was calculated with a Student’s t-test. Significance was accepted at the level of $P < 0.05$ (*), $P < 0.01$ (**) or $P < 0.001$ (***)

Materials and experimental procedures are detailed in the Extended Experimental Procedures.
Author Contributions
T.M. and O.T. designed the research, performed experiments and wrote the paper.
A.F. and T.F. performed polysome fractionation and ribosome-binding assay.
A.V. and D.M.S. carried out bioinformatics analysis.

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Accession Numbers
All data have been deposited at DDBJ: DRA003232 for Table S1, DRA003233 for Table S3, DRA003463 for Table S4 and DRA003234 for Table S6.
References


redundantly repress the Icos and Ox40 costimulator mRNAs and control follicular helper T cell differentiation. Immunity 38, 655-668.
Figure Legends

Figure 1. Identification of Reg1 target mRNAs

(A) SHAPE analysis of *in vitro* synthesized Il6 3’UTR (84-103) RNA.

(B and C) Schematic representation of Il6 3’UTR-(84-103) (B), and Il6 3’UTR and antisense morpholinos (C).

(D) HeLa cells were transfected with the Il6 3’UTR reporter, antisense morpholinos, and Reg1 expression plasmids. The luciferase activities are shown as relative values.

(E) Ctrl- and Il6-SL-MO were delivered into WT BMMs followed by stimulation with LPS. Cytokine mRNA (top) and protein (bottom) levels were analyzed by RT-qPCR and ELISA, respectively.

(F) WT BMMs delivered with antisense morpholinos were i.p. transfused into Il6−/− mice (6-week-old, n = 4). The mice were i.p. injected with LPS (0.5 mg/kg of BW). Cytokine levels in sera were measured by ELISA 24 h after transfusion.

(G) Schematic representation of λN-BoxB tethering system.

(H) Tef-off 293 cells were transfected with pTRE-II6CDS-5XBoxB and indicated expression plasmids. Total RNAs were prepared after Dox treatment, and Il6 levels were determined by Northern blot analysis.

(I) Immunoblot analysis of Reg1 in HeLa cells transfected with indicated expression plasmids.

(J) Correlation of ppm between control IP samples (x axis) and Reg1 IP samples (y axis). mRNAs associated with Reg1 in IL-1β-stimulated HeLa cells were determined by RIP-seq.

(K) Luciferase activity of HeLa cells transfected with reporter plasmids expressing 3’UTR of indicated genes and Reg1 expression plasmids.

Unless otherwise indicated, Data are mean ± SD (n = 3). See also Figures S1 and Table S1 and S2.
Figure 2. Identification of the consensus sequences of the Reg1 target mRNAs and their overlap with Roquin

(A and B) GSEA of overlap between Reg1 and Roquin target genes (RIP-seq based). ES plots are shown with genes ranked according to their Roquin (A) and Reg1 (B) RIP-seq p values. The right part shows the histogram for the max ES scores.

(C, G, J, K) Luciferase activity of HeLa cells transfected with indicated reporter, Reg1 and Roquin expression plasmids. Data are mean ± SD (n = 3).

(D) Phosphorimages of NuPAGE gels that resolved 5′-[32P]-labelled RNA–Flag-Reg1 IP. Lower panels show immunoblots for Reg1.

(E) Proportion of Reg1 binding sites obtained by the intersection of two biological HITS-CLIP replicates.

(F) Schematic representation of TM2D3 (95-120).

(H) SL enrichment in Reg1 bound CLIP tags in 3′UTRs compared to 1000 individual sequence permutations. The bar color represents the p value corresponding to the enrichment.

(I) Description of significantly enriched Reg1 bound SLs harboring a hairpin of 3 nucleotides with varying stem length (3, 5 or 7 nucleotides). Representation of SL sequence as Position Weight Matrix (PWM) logos. Logos are centered on the hairpin midpoint. Letter scale corresponds to relative nucleotide entropy.

See also Figure S2 and S3, and Table S3 and S4.

Figure 3. Reg1 localizes to ER and polysome fraction

(A, B) NIH3T3 cells expressing Flag-Reg1 were treated with arsenite and analyzed by confocal fluorescence microscopy.

(C) Immunoelectron microscopy showing the localization of Reg1 in NIH3T3 cells. The black arrows indicate Reg1 localized at surface of ER. Scale bar, 1 μm. mt, mitochondria.
(D) Subcellular fractions of Reg1 prepared from NIH3T3 cells were analyzed by Immunoblotting.

(E) The lysates from HeLa cells were fractionated by sucrose gradient. The expression of Reg1 and Roquin in the fractions was analyzed by Immunoblotting.

(F–H) Quantification of RNAs of non-polysome and polysome fractions. The lysates from HeLa cells transfected with indicated siRNA and stimulated with IL-1β were fractionated by sucrose gradient (F) and RNAs from the fractions were analyzed by RT-qPCR for the expression of indicated mRNAs (G, H). Data are mean ± SD (n = 3).

See also Figure S4.

Figure 4. Translationally active mRNAs are targeted by Reg1 via the stop codon

(A) WT and Reg1−/− macrophages were labelled with EU, stimulated with LPS and treated with anisomycin or CHX for 2 hr. EU-labeled Il6 and Ptgs2 mRNAs were captured by the Click-it Nascent RNA Capture Kit and measured by RT-qPCR.

(B and C) Tef-off 293 cells were co-transfected with pTRE-Il6 CDS-3’UTR and indicated expression plasmids. Cells were then treated with anisomycin or CHX, and RNAs were prepared after Dox treatment, and Il6 mRNA levels were determined by Northern blot analysis (B) and the autoradiographs were quantified (C).

(D) Schematic representation of SSL-inserted mRNAs.

(E) The SSL inhibited luciferase activity, but not its mRNA expression.

(F and G) Degradation of Il6, SSL-inserted Il6 and stop codon-deleted (∆STOP) Il6 by Reg1 was measured as in (B) (F), and the autoradiographs were quantified (G).

(H) Luciferase activity of HeLa cells transfected with the Il6 reporter lacking a stop codon and Reg1 expression plasmids.

(I and J) Schematic representation of deletion mutants of Il6 3’UTR (I). Degradation of Il6 mutant mRNAs was measured and quantified (J).

Unless otherwise indicated, Data are mean ± SD (n = 3).
Figure 5. Reg1 interacts with Ribosome proteins and UPF1

(A) Schematic representation of the iTRAQ workflow.
(B) Proteins enriched by iTRAQ-based proteomics identifying Reg1-associated proteins.
(C) Recombinant Reg1 co-precipitated with ribosome purified from HeLa cells was analyzed by SDS-PAGE.
(D and F) Immunoblot analysis of UPF1 co-precipitated with Flag-Reg1-WT and D141N (D) or its deletion mutants (F) in HeLa cells.
(E) Schematic representation of Reg1.
(G) Luciferase activity of HeLa cells transfected with indicated reporter and Reg1 mutant plasmids. Data are mean ± SD (n = 3).

See also Table S5.

Figure 6. UPF1 is essential for Reg1-mediated mRNA decay

(A and B) Reg1-mediated degradation of Il6 was measured in UPF1 knockdown cells (A) and the autoradiographs were quantified (B).
(C) Luciferase activity of HeLa cells transfected with UPF1, UPF2 or UPF3x siRNA, indicated reporter and Reg1 expression plasmids.
(D) Luciferase activity of UPF1 knockdown HeLa cells transfected with indicated reporter and the Roquin-1 expression plasmid.
(E) HeLa cells were transfected with the indicated siRNA and stimulated with IL-1β. RNA expression profiles were analyzed by RT-qPCR.
(F) Cells expressing Il6 were transfected with UPF1 siRNA and/or Flag-Reg1. Il6 co-precipitated with Reg1 were analyzed by RT-qPCR.
(G) Tethering Reg1 did not destabilize the luciferase-5BoxB under knockdown of UPF1.
Luciferase activity of UPF1 knockdown HeLa cells transfected with indicated reporter and expression plasmids.

(H) Degradation of Il6CDS-5BoxB mRNA under treatment with anisomycin was measured and quantified.

(I) Immunoblot analysis of UPF1 co-precipitated with Flag-Reg1 in HeLa cells under treatment with anisomycin.

(J-L) HeLa cells transfected with UPF1 siRNA were reconstituted with siRNA-resistant HA-UPF1-WTR or HA-UPF1-DEAA. Immunoblot analysis for the UPF1 reconstitution (J). Luciferase activity of reconstituted cells, after transfection of indicated reporter and Reg1 expression plasmids (K). RNA expression profiles of reconstituted cells stimulated with IL-1β (L).

Unless otherwise indicated, Data are mean ± SD (n = 3). See also Figure S6.

Figure 7. Roles of Reg1 and Roquin in the control of inflammatory mRNAs in response to LPS

(A-C) WT, Reg1+/− and RoquinSan/san BMMs were stimulated with LPS. RNA expression profiles were analyzed by RT-qPCR (A). Ptgs2 expression were analyzed by immunoblotting (B). The Protein/mRNA ratio of Ptgs2 is shown in (C).

(D) WT, Reg1+/−, RoquinSan/san, Reg1+/− and RoquinSan/san MEFs were stimulated with LPS and RNA expression profiles were analyzed by RT-qPCR.

(E) GSEA of overlap between Reg1 (left) or Roquin (right) target genes (RIP-seq based) and Reg1+/−; RoquinSan/San enrichment (transcriptome analysis based). An ES plot is shown with genes ranked according to their Reg1+/−; RoquinSan/San enrichment p value.

(F) GSEA of overlap between Reg1 (left) or Roquin (right) target genes (RIP-seq based) and Reg1+/− enrichment (transcriptome analysis based). An ES plot is shown with genes ranked according to their Reg1+/− enrichment p value.

(G) A proposed model of mRNA degradation by Reg1 and Roquin.
Unless otherwise indicated, Data are mean ± SD (n = 3). See also Figure S7 and Table S6.
## Table

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## Diagrams

### A

**Control (Empty)**
- 1) Benzonase treatment
- 2) Co-IP with anti-Flag Ab
- Labeling with iTRAQ-114
- Combine
- Quantify and identify by LC/MS/MS

**Flag-Reg1**
- 1) Benzonase treatment
- 2) Co-IP with anti-Flag Ab
- Labeling with iTRAQ-117
- Combine
- Quantify and identify by LC/MS/MS

### C

- Ribosome-binding Reg1
- Input Ribosome
- Input Reg1

### D

- Flag-Reg1-WT
- Flag-Reg1-D141N
- Myc-UPF1
- IB: Myc
- IB: Flag
- Benzonase

### E

**Mouse Reg1**
- Helix
- RNase
- Linker
- Zf

### F

- Flag-Reg1
- Myc-UPF1
- IB: Myc
- IB: Flag
- Benzonase

### G

- IL6 3’UTR-(1-403)
- Ptgs2 3’UTR-Full-(1-2100)
- Relative luciferase activity
### Figure A

(A) Relative mRNA expression of pro-inflammatory genes (IL6, TNF, PTGS2, NFKBIA) over time (0-8 hours) in WT, Roquin (San/San), and Reg1 (−/−) mice treated with LPS (0.5 mg/kg i.p.).

### Figure B

(B) Western blot images for Ptgs2 and Actin showing protein levels in WT, Roquin (San/San), and Reg1 (−/−) mice over time (0-8 hours) after LPS treatment.

### Figure C

(C) Bar graph showing protein/mRNA levels of Ptgs2 in WT and Reg1 (−/−) mice with or without Roquin (San/San) expression over time (0-8 hours) after LPS treatment.

### Figures D-F

(D) Relative mRNA expression of pro-inflammatory genes (IL6, TNF, PTGS2, NFKBIA) over time (0-8 hours) in WT, Reg1 (−/−), and Roquin (San/San) mice with or without the presence of Reg1 (−/−).

### Figures E-F

(E) Scatter plots showing enrichment scores (ES) for genes enriched in Reg1 RIP-seq compared to Roquin RIP-seq with a max. ES score of 0.3976 and p < 10^-4.

(F) Scatter plots showing enrichment scores (ES) for genes enriched in Reg1 RIP-seq compared to Roquin RIP-seq with a max. ES score of 0.2888 and p ~ 10^-4.

### Figures G

(G) Diagram illustrating the regulation of pro-inflammatory gene expression in acute and chronic phases of LPS-induced inflammation.

**Acute phase**
- Translationally Active mRNA
  - Ribosome, ER
  - Reg1
  - Stem-loop
  - mRNP
  - Translation termination
  - Recruitment of UPF1

**Chronic phase**
- Translationally Inactive mRNA
  - Stress granule, P body
  - Roquin
  - Stem-loop
  - mRNP
  - Deadenylase
  - Digestion of RNA

**Fine tuning of inflammatory gene expression**
Extended Experimental Procedures

Mice
Mice deficient in Reg1 have been described (Matsushita et al., 2009). Sanroque mice were kindly provided by Dr. Masakazu Hattori (Kyoto University). Il6−/− mice were kindly provided by Dr. Masao Matsuoka (Kyoto University). All animal experiments were done with the approval of the Animal Research Committee of the Institute for Virus Research, Kyoto University.

Cell culture and transfection
Primary MEFs were prepared from wild-type, Reg1−/−, Roquin san/san or Reg1−/−/Roquin san/san mouse embryos at embryonic day 13.5. Dicer-ablated MEFs were provided by Stephen N. Jones (Mudhasani et al., 2008). MEFs, NIH3T3 cells, RAW cells, HeLa cells and HEK293 cells were maintained in DMEM (Nacalai Tesque) supplemented with 10% (vol/vol) FBS and 50 μM β-mercaptoethanol (Invitrogen). Tet-off HEK293 cells were purchased from Clontech and maintained in α-MEM (Nacalai Tesque) supplemented with 10% (vol/vol) FBS (Clontech), 50 μM β-mercaptoethanol and 100 μg/mL G418 (Nacalai Tesque). Flp-In 293 T-REx cells (Invitrogen) were grown in DMEM high glucose with 10% (v/v) FBS and 2 mM L-glutamine. Cells were transfected through the use of Lipofectamine 2000, Lipofectamine 3000, Lipofectamine LTX (Invitrogen), the Neon transfection system (Invitrogen) or the Nucleofector system (Lonza) according to the manufacturer’s recommendations.

For siRNA-mediated knockdown, cells were transfected through the use of Lipofectamine RNAiMAX (Invitrogen), MISSION siRNA Transfection Reagent (Sigma) or the Neon transfection system (Invitrogen). The siRNAs used in this study were synthesized by Ambion and the following siRNA target sequences were used: hUPF1, 5′-GAUGCAGUUCCGCUCUCAUUdTdT-3’; mUPF1,
Peritoneal macrophages were prepared from mice 3 d after intraperitoneal injection of 4% (vol/vol) thioglycollate medium (2 ml) (Sigma) and were maintained in RPMI-1640 medium (Nacalai Tesque) supplemented with 10% (vol/vol) FBS and 50 μM β-mercaptoethanol.

For preparation of bone marrow–derived macrophages (BMMs), bone marrow cells were isolated from wild-type, Reg1−/−, or Roquinsan/san mice and cultured in macrophage growth medium (RPMI-1640 medium supplemented with 10% (vol/vol) FBS, 50 μM β-mercaptoethanol, 100 U/ml of penicillin, 100 μg/ml of streptomycin and 20 ng/ml of macrophage colony-stimulating factor). After 5 days, cells were washed once and cultivated for 2 days with macrophage growth medium, then cells were collected for further analysis.

**Plasmid construction and reagents**

The cDNAs of mouse Reg1 (Zc3h12a), the D141N mutant and mouse Il6 have been
described (Matsushita et al., 2009). The cDNAs of Reg1 were ligated to the vector pFlag-CMV2 (SIGMA) and pcDNA3.1(+) (Invitrogen) for mammalian expression, pMRX-ires-puro (Cell Biolabs) for retrovirus production, pGEX-6P1 (GE Healthcare) for *E. coli* expression and pTNT vector (Promega) for protein production. The cDNAs of Il6 CDS-3′ UTR and β-globin CDS-Reg1 3′ UTR were inserted in pTREtight vector (Clontech). The 3′ UTR cDNAs of Il6 were inserted downstream of the T7 promoter of pBluescript II KS (−) (Stratagene). The 3′ UTR cDNAs of a set of genes were inserted in the pGL3-promoter (Promega). pENTR4 constructs were generated by PCR amplification of the Reg1 coding sequences (CDS) from cDNA followed by restriction digest and ligation into the pENTR4 (Invitrogen) backbone, which were further recombined into the pFRT/TO/FLAG/HA-DEST destination vector (Baltz et al., 2012) using GATEWAY LR recombinase (Invitrogen) according to manufacturer's protocol.

TOPuro-hCaf1a-AA-mycSG, pcDNA3-Flag-hDcp2-AA, pLNCX2-EGFP-mRoquin, and pCI-neo-Flag-mRoquin were kindly provided by Georg Stoecklin (German Cancer Research Center). pSR-myc-hUPF1-WT, siRNA-resistant pSR-HA-hUPF1-WT(R) and pSR-HA-hUPF1-D648A/E649A (DEAA) (R) were kindly provided by Akio Yamashita (Yokohama City University).

LPS from *Salmonella minnesota* and puromycin were from Invivogen. Recombinant cytokines were from R&D Systems. Anisomycin, cycloheximide and actinomycin D were from Sigma; Harringtonine was from LKT Laboratories.

**In vitro RNA cleavage assay**

In vitro RNA cleavage assay has been previously described (Matsushita et al., 2009). Briefly, Recombinant Reg1 protein (Matsushita et al., 2009) and in vitro transcribed 5′-[32P]-labelled RNAs were mixed in cleavage buffer (25 mM HEPES, 50 mM potassium acetate, 5 mM DTT, 5 mM magnesium acetate and 0.2 U/ml RNasin (Promega)). The cleaved RNA was purified with Trizol (Invitrogen) and analysed by
denaturing 6% polyacrylamide-TBE-urea gels (Invitrogen) and autoradiography.

For generation of the circular RNA, in vitro transcribed 5′-[32P]-labelled mIl6 3′ UTR (1-403) RNA was circularised by T4 RNA Ligase 1 (ssRNA Ligase) (NEB) in ligation buffer (50 mM Tris-HCl, pH 7.5, 10 mM MgCl2, 1 mM DTT, 10% PEG8000, 50 μM ATP and 0.2 U/ml RNasin (Promega)) at 25°C for 2 h. The RNA product was subjected to in vitro RNA cleavage assay as described above.

RNA motif exploring by selective 2′-hydroxyl acylation analyzed by primer extension (SHAPE) chemistry and computational analysis

RNA SHAPE chemistry has been described (Wilkinson et al., 2006). Briefly, DNA templates for mIl6 3′ UTR (84-103), mPtgs2 3′ UTR (1204-1237) or stem-loop structure were inserted in the context of flanking 5′ and 3′ structure cassette (Wilkinson et al., 2006), which was inserted downstream of the T7 promoter of pBluescript II KS (−) (Stratagene), and in vitro transcribed by T7 RNA polymerase (MEGAscript T7 Transcription Kit, Invitrogen). The in vitro transcribed RNAs were refolded in RNA folding buffer (100 mM Tris-HCl, pH 8.0, 100 mM NaCl and 6 mM MgCl2) in a final volume of 18 μl. After folding, RNAs were modified in the presence of 6.5 mM N-methylisatoic anhydride (NMIA) (M-25; Invitrogen) and incubated at 20°C for 4 h. After modification, RNAs were subjected to reverse transcription (ReverTra Ace, Toyobo) using a 5′-[32P]-labelled DNA primer (5′-GAACCGGACCGAAGCCCG-3′) specific to the 3′ structure cassette. For G or A sequencing experiment, primer extensions were performed in the presence of cytosine or thymidine dideoxy terminating nucleotides, respectively. The cDNA fragments are resolved on denaturing 8% polyacrylamide-TBE-urea sequencing gels followed by autoradiography. Computational prediction of the RNA secondary structures was performed using the RNAfold from the Vienna package (Lorenz et al., 2011).
Tethered Function Assays

Reg1 was artificially recruited to reporter mRNAs by the BoxB-λN-peptide tethering system (Mishima et al., 2012). The 22 amino acid RNA-binding domain of λN peptide was fused with the N terminus of Reg1 and five 19 nt λN binding sequences (5XBoxB) were tandemly inserted in a downstream of Il6 coding sequence (CDS) or luciferase CDS. Tet-off 293 cells and HeLa cells were transfected with the 5XBoxB reporter plasmids, together with expression plasmid for Flag-λN-Reg1 or empty (control) plasmid followed by luciferase assay, mRNA decay assay and RNA IP.

RNA immunoprecipitations (RIP)-sequencing analysis

RNA immunoprecipitation was performed using a RIP-Assay Kit (RN1001; Medical & Biological Laboratories) according to the manufacturer's protocol with some modifications. Briefly, HeLa cells were transfected with expression vector for Flag-tagged Reg1-D141N mutant, Roquin-1 or empty (control) vector and stimulated with IL-1β (10 ng/mL) for 0 or 2 h. RNA-protein complexes from lysate of HeLa cells were immunoprecipitated with anti-Flag antibody (F3165; Sigma) bound to protein G magnetic beads and RNAs were extracted from the beads. The quality of RNA was analyzed by the Bioanalyzer Nano 6000 chip (Agilent Technologies). Only samples with an RNA Integrity Number (RIN) greater than 9.0 were used for further experiments. RNA library was prepared using Small RNA Sample Prep kit v1.0 (Illumina Inc.) and sequenced on a HiSeq 2000 system (Illumina) also according to the manufacturer's instructions. Fourteen and 0.6 million RNA reads were obtained for Reg1 and 16.1 million for control. The resulting set of trimmed reads were then mapped against the human genome (hg19; NCBI). Analysis of enrichment of mapped reads in the Reg1 RIP samples vs control IP samples was performed using the R package DESeq (Anders and Huber, 2010), as follows. The count of mapped reads to each gene was used as input for the method. The coverage of each sample and the dispersions were estimated from the
read count data using the functions provided in the package. Finally, differential tag counts for each gene between the RIP and control sample for each of the two time points were estimated based on the negative binomial distribution. Genes with an unadjusted p value for enrichment in the RIP samples ≤ 1e-3 were regarded as candidate Reg1 targets.

**Analysis of overlap with between Reg1 targets and Roquin targets.**

The overlap between the targets of Reg1 and those of Roquin was evaluated using three measures. First, we employed the Gene Set Enrichment Analysis (GSEA) methodology, as described (Subramanian et al., 2005). In this case, we defined a set of genes S as the Reg1 targets based on their enrichment in RIP-seq samples at 0h, 2h, and both time points combined. On the other hand, a ranked list L was obtained by sorting the genome-wide set of genes by their enrichment in the Roquin RIP-seq samples of the same time points. The distribution of S in L was evaluated by calculating an Enrichment Score (ES) and the estimation of a significance level for ES by 10,000 random permutations, as described (Subramanian et al., 2005). The P value of enrichment of S in the top of L was estimated as the fraction of permuted samples resulting in a higher ES than the actual data.

The statistical significance of overlap between Reg1 targets and Roquin targets was also evaluated using the Kolmogorov-Smirnov statistical test, which is closely related to the GSEA approach, and the hypergeometric distribution. In the latter case, Roquin targets were defined as the 52 genes having a p value of enrichment in the RIP sample smaller than 0.01 at either of the time points, and Reg1 targets as 68 genes having a p value of enrichment < 0.001 at either of the time points. The number of overlapping targets of both time points were compared to the expected number according to the hypergeometric distribution. Both the Kolmogorov-Smirnov and the hypergeometric test were performed using the R language. Candidate mouse Roquin targets were
obtained from (Leppek et al., 2013). The 92 human homologs of the mouse targets were defined using human-mouse homology data as available from the Mouse Genome Database (MGD) at the Mouse Genome Informatics website, The Jackson Laboratory (http://www.informatics.jax.org; April 2014).

**Generation of NIH3T3 cells stably expressing Flag-tagged Reg1**

NIH3T3 cells were transduced with retroviral supernatant for 12 h. Retroviral supernatants were produced by PlatE packaging cells (Cell Biolabs) transfected with retroviral expression vectors. Supernatants were collected 48 h after transfection, were filtered through 0.22-μm filters and were used for infection. After transduction, cells were washed and resuspended for 12 h in maintenance media. Cells were washed once and cultured in maintenance media with 2 μg/ml puromycin. Stable clones were obtained by selection in the presence of puromycin (2 μg/mL) and analyzed by Western blotting for the expression of Flag-tagged Reg1 using the anti-Flag antibody M2 (F3165; Sigma).

**Fluorescence microscopy**

For immunofluorescence staining, NIH3T3 cells stably-expressing Flag-tagged Reg1 were used because of large cell cytoplasm in this cell type. In addition, HeLa cells transiently-expressing Flag-tagged Reg1 were also used. The cells were fixed in 3% paraformaldehyde in PBS for 15 min, incubated with 50 mM NH₄Cl in PBS for 10 min, permeabilized in 0.5% Triton X-100 and 0.1% gelatin in PBS for 10 min, washed three times in 0.1% gelatin in PBS and blocked in 0.1% gelatin and 2% goat-serum (X0907; Dako) in PBS for 30 min at room temperature. Primary antibodies to Flag (F3165 or F7425; Sigma), α/β-Tubulin (#2148; Cell Signaling Technology), Calnexin (ab22595; Abcam), PDI (#3501; Cell Signaling Technology), RPL7A (15340-1-AP; Proteintech Group), Dcp1a (ab47811; Abcam) or TIA1 (ab40693; Abcam) were used for staining in
combination with secondary antibodies conjugated to Alexa 488 (A11017 or A11070; Invitrogen) or Alexa 568 (A11019 or A21069; Invitrogen). Images were captured on a Leica TCS SPE confocal microscopes (Leica) and an All-in-one Type Fluorescence Microscope (BZ-9000; Keyence) and analyzed with the LAS-AF software (Leica) and BZ Analyzer Software (Keyence), respectively.

For RNA fluorescence in situ hybridization (RNA FISH) combined with immunofluorescence (IF) staining, HeLa cells were stimulated with IL-1β (10 ng/ml) for 4 h and were subjected to RNA FISH. RNA FISH was performed using the RNAscope Multiplex Fluorescent Assay (Advanced Cell Diagnostics) according to the manufacturer's protocol with some modifications. Briefly, cells were fixed in 4% paraformaldehyde in PBS for 20 min, serially dehydrated and rehydrated for 5 min each in graded alcohol (50%, 70% and 100% EtOH), treated with protease for 10 min at room temperature and hybridized with RNAscope probes at 40°C. After the washing step, cells were blocked in 1% (wt/vol) Blocking Reagent (11096176001; Roche) in PBS for 1 h at room temperature. Primary antibodies to Dcp1a (ab47811; Abcam) were used for staining in combination with secondary antibodies conjugated to Alexa 568 (A21069; Invitrogen). Images were captured on a Leica TCS SPE confocal microscopes (Leica) and analyzed with the LAS-AF software (Leica).

**Immunoelectron microscopy**

Immunoelectron microscopy has been described (Matsunaga et al., 2009). Briefly, NIH3T3 cells stably-expressing Flag-tagged Reg1 were fixed with 4% paraformaldehyde for 30 min in 0.1 M sodium-phosphate buffer (pH 7.4) and washed for 5 min three times in the sodium-phosphate buffer. Cells were permeabilized and blocked for 30 min with 0.2% saponin, 10% BSA, 10% normal goat serum and 0.1% cold-water fish gelatin in the sodium-phosphate buffer. Cells were stained with anti-Flag mouse monoclonal antibody (F3165; Sigma) overnight at 4°C, washed for 10 min six
times in the sodium-phosphate buffer containing 0.1% saponin, then stained for 2 h at room temperature with an anti-mouse IgG conjugated to 1.4 nm gold particle (Nanogold Fab’ fragment of goat anti-mouse IgG) (2002; Nanoprobes), washing for 10 min five times in the sodium-phosphate buffer containing 0.1% saponin. Electron micrographs were obtained with a JEM-1011 transmission electron microscope (JEOL).

**iTRAQ labeling and mass spectrometry**

iTRAQ labeling was performed using an iTRAQ reagent kit (Applied Biosystems) according to the manufacturer's protocol with some modifications. Briefly, cell lysates from RAW cells stably expressing Flag-tagged Reg1 and HeLa cells transiently-expressing Flag-tagged Reg1 together with their controls (Empty) were immunoprecipitated with anti-Flag Ab in the presence of Benzonase (50 U/mL) (Novagen). The immunoprecipitated proteins (5 µg each) were reduced, alkylated, digested, labeled with iTRAQ reagents and cleaned up with a cation exchange chromatography according to the manufacturer's (Applied Biosystems) protocol with some modifications. Control or Reg1 immunoprecipitates was labeled with iTRAQ Reagent 114 or 117, respectively. The iTRAQ reagent-labeled sample was analyzed by LC-MS/MS (Orbitrap LC-MS; Thermo Fisher Scientific). Proteins were identified by database searching using Mascot Daemon (Matrix Science). Based on an 80% confidence level (< 20% 117/114 variability), cutoff values of 1.2-fold for up-regulated proteins were used to define candidate proteins as Reg1-associated proteins.

**Sucrose gradient centrifugation (polysome profiles)**

HeLa cells were lysed in polysome buffer (20 mM HEPES-KOH [pH 7.5], 100 mM KCl, 5 mM MgCl₂, 0.25% (vol/vol) Nonidet P-40, 10 µg/ml cycloheximide, 100 units/ml RNase inhibitor, protease inhibitor cocktail). Lysates were loaded on top of a linear 15%–60% sucrose gradient (15%–60% sucrose, 20 mM HEPES-KOH [pH 7.5],
100 mM KCl, 5 mM MgCl₂, 10 µg/ml cycloheximide, 100 units/ml RNase inhibitor, protease inhibitor cocktail). After ultracentrifugation at 38,000 rpm for 2.5 h at 4°C in a HITACHI P40ST rotor, fractions were collected from the top of the gradient and subjected to UV-densitometric analysis. The absorbance profiles of the gradients were determined at 254 nm. For disassociation of ribosome and polysome, EDTA was added to Mg²⁺-free polysome buffer and 15%–60% sucrose gradient at concentrations of 50 mM and 20 mM, respectively. For protein analysis, each fraction was subjected to TCA precipitation and SDS-PAGE. Immunoblotting was performed using the indicated antibodies. For RNA analysis, each fraction was subjected to High Pure RNA Isolation Kit (Roche) and RT-qPCR.

**Immunoblot analysis**

Whole-cell extracts were prepared in lysis buffer (1% (vol/vol) Nonidet P-40, 0.1% (wt/vol) SDS, 1% (wt/vol) sodium deoxycholate, 150 mM NaCl, 20 mM Tris-HCl, pH 8.0, 10 mM EDTA and Complete Mini Protease Inhibitor Cocktail without EDTA (Roche)). The following antibodies were used for immunoblot analysis: antibody to Flag (F3165 and F7425; Sigma), c-Myc (M4439 and C3956; Sigma), α/β-Tubulin (#2148; Cell Signaling Technology), Calnexin (ab22595; Abcam), PDI (#3501; Cell Signaling Technology), Roquin-1 (A300-514A; Bethyl Laboratories, Inc.), Roquin-2 (ab99090; Abcam), Staufen1 (STAU1) (A303-956A; Bethyl Laboratories, Inc.), UPF2 (ab153830; Abcam), UPF3 (ab83249; Abcam), Dep1a (ab47811; Abcam), TIA1 (ab40693; Abcam), RPL7A (15340-1-AP; Proteintech Group), RPS4X (14799-1-AP; Proteintech Group), RPL7 (ab72550; Abcam), Ptgs2 (ab15191; Abcam) and β-actin (sc-1615; Santa Cruz). Rabbit UPF1 antibody was kindly provided by Akio Yamashita (Yokohama City University). Rabbit Reg1 antibody has been described (Iwasaki et al., 2011). Luminescence was detected with a luminescent image analyzer (ImageQuant LAS 4000; GE Healthcare).
Subcellular fractionation

Subcellular fractionation was performed using an Endoplasmic Reticulum Isolation Kit (Sigma) following the manufacturer’s instructions. Briefly, NIH3T3 cells were suspended in hypotonic extraction buffer (10 mM HEPES, pH 7.8, 25 mM potassium chloride and 1 mM EGTA) to allow the cells to swell. Cells were adjusted to isotonic conditions by the addition of isotonic extraction buffer (10 mM HEPES, pH 7.8, 250 mM sucrose, 25 mM potassium chloride and 1 mM EGTA) and homogenized by the Dounce homogenizer. The total lysate was centrifuged at 1,000 x g for 10 min at 4°C, yielding a pellet consisting mainly of nuclei and unbroken cells. The supernatant was centrifuged at 12,000 x g for 15 min at 4°C, yielding a pellet consisting mainly of mitochondria. The supernatant was ultracentrifuged at 100,000 x g for 1 h at 4°C, yielding a pellet consisting mainly of microsomal fraction (cellular membranes, SER and RER). The supernatant from this 100,000 x g spin consists mostly of cytosol. The pellet consisting of microsomal fraction was resuspended in the isotonic extraction buffer and applied to an Optiprep gradient and ultracentrifuged at 150,000 x g for 3 h at 4°C to obtain SER and RER fractions. Each fraction was collected and analyzed by Western blot with the indicated antibodies.

Recombinant Reg1 proteins

Flag-tagged Reg1 protein and its mutant protein were translated in vitro in the presence of [35S]-methionine with the TNT-coupled reticulocyte lysate system (Promega) following the manufacturer’s instructions with some modifications. After in vitro translation, lysates were treated with Benzonase (Novagen) and [35S]-labeled proteins were affinity purified using anti-Flag antibody affinity resin (Sigma) and eluted with a 3 x Flag peptide (Sigma). The [35S]-labeled proteins were characterized with quantitative autoradiography.
**Ribosome-binding assay**

Ribosomes were purified from HeLa cells by sedimentation through a sucrose cushion (Zinzalla et al., 2011). Recombinant $[^{35}S]$-radioactive Reg1 protein was incubated in ribosome binding buffer (10 mM Tris-HCl, pH 7.6, 100 mM KCl, 10 mM MgCl$_2$, 1 mM DTT, 0.1 mg/mL BSA) for 30 min at 4°C in the simultaneous absence or presence of the highly-purified ribosome (5 µg). Subsequently the mixture was loaded on a 10% sucrose cushion (10 mM Tris-HCl, pH 7.6, 100 mM KCl, 10 mM MgCl$_2$, 10% (wt/vol) Sucrose) and ultracentrifuged for 20 min at 300,000 x g at 4°C to pellet the ribosome and ribosome-associated protein. The pellets were analyzed by SDS-PAGE and quantitative autoradiography.

**Quantitative PCR analysis**

Total RNA was isolated using ISOGEN II (Wako) or Trizol (Invitrogen) and reverse-transcribed using ReverTra Ace (Toyobo) or MicroRNA RT Kit (Applied Biosystems) according to the manufacturer’s instructions. For quantitative PCR, cDNA fragments were amplified through the use of Thunderbird Probe qPCR Mix (Toyobo) or TaqMan® MicroRNA Assays (Applied Biosystems); TaqMan probes for mouse Il6, mouse β-Actin, mouse 18S rRNA, human IL6, human 18S rRNA and human β-ACTIN were from Applied Biosystems. Fluorescence was detected with a StepOne Real-Time PCR System (Applied Biosystems). For analysis of the induction mRNA in response to LPS stimulation, the abundance of mRNA of each expressed gene was normalized to that of 18S rRNA or β-Actin.

**Protein IP**

HeLa cells and Tet-off 293 cells transiently-expressing Flag-tagged Reg1 and RAW cells and Jurkat cells stably-expressing Flag-tagged Reg1 were lysed in lysis buffer
(0.5% (vol/vol) Nonidet P-40, 150 mM NaCl, 10 mM Tris-HCl, pH 8.0 and Complete
Mini Protease Inhibitor Cocktail without EDTA (Roche)). Anti-Flag antibody (F3165;
Sigma) bound to protein G magnetic beads (Invitrogen) was incubated for 3 h at 4ºC
with lysates. Beads were washed three times with lysis buffer and suspended in SDS
sample buffer (50 mM Tris-HCl, pH 6.8, 2% (wt/vol) SDS, 5% (vol/vol)
β-mercaptoethanol, 10% (vol/vol) glycerol and bromophenol blue). Samples were
boiled for 5 min at 95ºC and analyzed by Western blot with the indicated antibodies.

**RNA IP**

HeLa cells and Tet-off 293T cells transfected with the indicated siRNA and expression
plasmids and RAW cells stably expressing Flag-tagged Reg1 were lysed in RNA IP
lysis buffer (0.5% (vol/vol) Nonidet P-40, 150 mM NaCl, 20 mM Tris-HCl, pH 7.5,
Complete Mini Protease Inhibitor Cocktail without EDTA (Roche) and 0.2 U/ml RNasin
(Promega)). Anti-Flag antibody (F3165; Sigma) bound to protein G magnetic beads
(Invitrogen) was incubated for 3 h at 4ºC with lysates and beads were washed three
times with RNA IP lysis buffer. RNAs were eluted from the beads using Trizol reagent
(Invitrogen) according to the manufacturer’s instructions and analyzed by RT-qPCR.
Proteins were eluted from the beads using SDS sample buffer (50 mM Tris-HCl, pH 6.8,
2% (wt/vol) SDS, 5% (vol/vol) β-mercaptoethanol, 10% (vol/vol) glycerol and
bromophenol blue) and analyzed by Western blot with the indicated antibodies.

**Morpholino delivery and ELISA for sera and culture supernatants**

The antisense morpholino oligonucleotides used in this study were synthesized by Gene
Tools, LLC and the following 25-mer morpholinos were used: Il6-SL-MO,
5’-CATTCATATTGTCAGTTCTTCGTAG-3’; and Ctrl-MO as the standard control
oligo from Gene Tools, LLC (SC100), 5’-CCTCTTACCTCAGTTACAATTTATA-3’.
For delivery of antisense morpholino oligonucleotides to HeLa cells or bone marrow
macrophages (BMMs) from wild-type mice, morpholinos were added to the medium at a final concentration of 2 μM in the presence of 6 μM Endo-Porter (Gene Tools, LLC) according to the manufacturer’s protocol, and cells were incubated for 24 h and used in subsequent experiments. HeLa cells were used in luciferase assay as described above.

For measurement of Il6 and Tnf concentrations in culture supernatants from BMMs, BMMs were stimulated with LPS (100 ng/mL) and cytokine levels in culture supernatants were measured by cytokine-specific ELISA kits (R&D systems) following manufacturer’s protocols. Cytokine mRNA levels in BMMs were analyzed by RT-qPCR as described above.

For measurement of Il6 and Tnf concentrations in sera from Il6–/– mice, BMMs (5 x 10⁶ cells) were intraperitoneally (i.p.) transfused into Il6–/– mice (6-week-old, n = 4). The mice were injected with LPS (0.5 mg/kg of body weight, i.p.) 24 h after transfusion, and cytokine levels in sera were measured by cytokine-specific ELISA kits (R&D systems) following manufacturer’s protocols.

**HITS-CLIP**

Flp-In 293 T-REx cell lines stably expressing FLAG/HA-tagged Reg1 protein were generated by co-transfection of pFRT/TO/FLAG/HA constructs with pOG44 (Invitrogen). Cells were selected by adding 15 μg/ml blasticidin and 100 μg/ml hygromycin (Invivogen). Expression of epitope-tagged proteins was induced by addition of 1 μg/ml doxycyclin. The expression of FLAG/HA tagged Reg1 protein was assessed by Western analysis using mouse anti-HA.11 monoclonal antibody (Covance). HITS-CLIP was performed essentially as described (Hafner et al., 2010). Briefly, UV-irradiated cells were lysed in NP-40 lysis buffer (50 mM HEPES-KOH at pH 7.4, 150 mM KCl, 2 mM EDTA, 0.5% (v/v) NP40, 0.5 mM DTT, complete EDTA-free protease inhibitor cocktail). After treatment with RNaseT1 (Fermentas) at final concentration of 0.5 unit/μl, immunoprecipitation was carried out with Flag magnetic
beads (SIGMA) from HEK293 cell extracts for 1 h at 4°C. Following additional
digestion by RNase T1 (Fermentas) at final concentration of 0.5 unit/µl, beads were
incubated with calf intestinal phosphatase (NEB) and RNA fragments were
radioactively end-labeled using T4 polynucleotide kinase (Fermentas). The crosslinked
protein-RNA complexes were resolved on a 4-12% NuPAGE gel (Invitrogen). The
SDS-PAGE gel was transferred to a nitrocellulose membrane (Whatman) and the
protein-RNA complex migrating at an expected molecular weight was excised. RNA
was isolated by Proteinase K (Roche) treatment and phenol-chloroform extraction,
ligated to 3’ adapter and 5’ adapter, reverse transcribed and PCR-amplified. The
amplified cDNA was sequenced on a HighSeq2000 (Illumina) with a 1x51 nt cycle.

Reg1 CLIP library processing

Reads from the Reg1 CLIP library were stripped of the adaptor sequence stepwise using
Cutadapt, keeping the randomized nucleotides (Martin, 2011) (3’ Adapter, NN-TGGAATTCTCGGTGCCAAGG; 5’ Adapter, GTTCAGAGTTCTACAGTCCGATC-NN). Reads that were less than 24
nucleotides in length or contained an ambiguous nucleotide were discarded. Reads were
collapsed and then the randomized adapter nucleotides were trimmed using Flexbar
(Dodt et al., 2012). The remaining reads were aligned to the human genome (hg19),
with up to one mismatch allowed and ten alignment locations, with Bowtie version
0.12.7 (Langmead et al., 2009). Grouping of the reads was done using PARalyzer
(Corcoran et al., 2011) to retain only those that mapped to a single genomic location for
the minimum number of mismatches. Unique reads were grouped with a minimum of
one nucleotide of overlap. Annotation of the resulting groups was performed as
described (Ascano et al., 2012). The two libraries were intersected on group level with
at least one nucleotide overlap. Full group level data are provided in Table S4.
Stem loop enrichment and stem loop sequence analyses

Group sequences were globally folded using RNAfold from the Vienna package (Lorenz et al., 2011) with default setting. Resulting stem loops have been quantified and compared against the number of retrieved stem loops after 1000 individual dinucleotide shuffles (using Dishuffle (Clote et al., 2005)) of the corresponding group sequences. Stem loop Postion weight matrixes were obtained using the web tool enoLOGOS (Workman et al., 2005), using stem loop sequences as input. RNA secondary structure and sequence motifs were predicted using Predict Motif (Rabani et al., 2008), using the entire Reg1 bound group sequence for the corresponding stem loop as input.
References for Extended Experimental Procedures


Supplemental Figure Legends

Figure S1. Identification of novel Reg1 target molecules, related to Figure 1

(A) Reg1 endonucleolytically cleaved the Il6 3'UTR (1-403) \textit{in vitro} cleavage assay. Indicated circular or linear Il6 3'UTR RNAs were incubated with increasing amounts of recombinant Reg1 proteins.

(B) Coomassie Brilliant Blue-stained SDS gel showing purified recombinant WT or D141N mutant Reg1 proteins.

(C) Reg1 cleaved the (84-103) SL of Il6-3'UTR \textit{in vitro} cleavage assay. Indicated Il6 RNAs were incubated with increasing amounts of recombinant Reg1 proteins.

(D) Luciferase activity of HeLa cells transfected with reporter plasmids expressing 5XBoxB and Il6 3'UTR, together with indicated expression plasmids. The gene encoding Renilla luciferase was transfected simultaneously as an internal control. Data are mean ± SD.

(E) MEFs from \textit{Reg1}^{-/-} and WT mice were stimulated with LPS (100 ng/mL) and miRNA expression levels were analyzed by RT-qPCR. Data are mean ± SD (n = 3).

(F) Structural models of WT and D141N mutant Reg1 PIN domains. Cartoon representations of homology models of the PIN domain (residues 132-295) of WT and D141N mouse Reg1 after 10 ns of explicit water MD. Regions colored red are predicted to interact with RNA.

(G) Luciferase activity of HeLa cells transfected with indicated reporter and indicated Roquin expression plasmids. Roquin-NT is an N-terminal fragment (amino acids [aa] 2–441), which is responsible for RNA-binding. Data are mean ± SD.

Figure S2. Identification of Reg1-recognition motives, related to Figure 1

(A, D, E, I) Luciferase activity of HeLa cells transfected with indicated reporter and expression plasmids. Data are mean ± SD.

(B, H) Schematic representation of Reg1-recognition sequences in mouse \textit{Nfkbiz} (B)
and Ptgs2 (H) together with an alignment of 5 mammalian sequences.

(C, F, J,) Schematic representation of Reg1-recognition stem-loop structures in Nfkbiz (C), Ptgs2 (F), TNE-CDE (J).

(G) SHAPE analysis of in vitro synthesized mPtgs2 3′UTR (1204-1237) RNA.

**Figure S3. Reg1 recognizes stem-loop motifs in 3'UTRs, related to Figure 2**

(A, B) Schematic representation of TNF-CDE (A) and an artificial stem-loop (B) structures.

(C) SHAPE analysis of in vitro synthesized a “UAU-loop” stem-loop RNA.

(D, E) Luciferase activity of HeLa cells transfected with indicated reporter and Reg1 expression plasmids. Data are mean ± SD.

(F–H) mRNAs associated with Reg1 in HeLa cells transfected with indicated expression plasmids were recovered by RNA IP and analyzed RT-qPCR. Data are mean ± SD. Lower panels show immunoblots probed with an anti-Flag antibody.

**Figure S4. Reg1 and Roquin degrade mRNAs via distinct mechanisms, related to Figure 3**

(A) Top: Degradation of Il6CDS-3′UTR mRNA under co-transfection with Caf1a-AA or Dcp2-AA was measured as in Figure 4B. Bottom: Quantification of the autoradiographs. Data are mean ± SD.

(B and C) Immunofluorescent microscopy showing the localization of Roquin to PB and SG. NIH3T3 (B) and HeLa (C) cells transiently-expressing EGFP-tagged Roquin were treated with arsenite for 1 h and analyzed by confocal fluorescence microscopy.

(D) Immunoblot analysis of endogenous and Flag-Reg1 expression in Flag-Reg1 stably expressing NIH3T3 cells. The levels of Reg1 detected by ant-Reg1 Ab was quantified and shown.

(E) Immunofluorescent microscopy showing the localization of Reg1 to the cytoplasm
and ER, but not to PB and SG. HeLa cells transiently-expressing Flag-Reg1 were analyzed by confocal fluorescence microscopy.

(F) Fluorescence *in situ* hybridization of *Ptgs2* and fluorescence immunostaining of Dcp1a (PB marker) in HeLa cells transfected with siRNA specific for Reg1 or Roquin-1/2 and stimulated with IL-1β for 4 h.

(G) Quantification of *Ptgs2* mRNAs in Figure S4F. Data are mean ± SD.

(H) Quantification of *Ptgs2* mRNAs localized in PB (%) in Figure S4F. Data are mean ± SD.

**Figure S5. Treatments with protein synthesis inhibitors have no effect on the localization of Reg1, related to Figure 4**

(A) NIH3T3 cells expressing Flag-Reg1 were treated with anisomycin or CHX for 2 h and analyzed by confocal fluorescence microscopy.

(B) Immunoblot analysis of Reg1 transiently-expressed in Tet-off 293 cells under treatment with anisomycin or CHX as in Figure 4B and 4C.

(C) IRES-dependent translation initiation had no effect on Reg1-mediated RNA decay. Degradation of IRES-*Il6* mRNA was inserted in upstream of *Il6* CDS, was measured as in Figure 4B.

(D) Quantification of the autoradiographs in Figure S5C. Data are mean ± SD.

**Figure S6. Essential roles of UPF1 in Reg1-mediated mRNA decay, related to Figure 6.**

(A) Immunoblot analysis of UPF1 and Reg1 in RAW264.7 macrophages transfected with indicated siRNAs.

(B) Luciferase activity of RAW264.7 macrophages transfected with indicated siRNA, reporter and Reg1 expression plasmids. Data are mean ± SD.

(C) Luciferase activity of RAW264.7 macrophages transfected with indicated siRNA,
reporter and Roquin expression plasmids. Data are mean ± SD.

(D) RAW264.7 macrophages were transfected with the indicated siRNAs and stimulated with LPS. RNA expression profiles were analyzed by RT-qPCR. Data are mean ± SD.

(E) Immunoblot analysis of Reg1 in HeLa cells transfected with indicated expression plasmids.

(F) mRNAs associated with Reg1 in Tet-off 293 cells treated with anisomycin were recovered by RNA IP and analyzed RT-qPCR. Data are mean ± SD.

(G) Immunoblot analysis of UPF1 and its helicase dead mutant (DEAA) co-immunoprecipitated with Flag-Reg1-WT transiently-expressed in HeLa cells.

Figure S7. Inflammatory gene expression in cells defective in the Reg1 and Roquin to TNF stimulation, related to Figure 7

(A) HeLa cells transiently-expressing Flag-Reg1-D141N were analyzed by confocal fluorescence microscopy.

(B) Luciferase activity of HeLa cells transfected with indicated reporter and Reg1 and Roquin expression plasmids. Data are mean ± SD.

(C) The ratio of non-polysome and polysome RNA to total RNA in Figure 3H.

(D) MEFs from Reg1−/−, RoquinSan/san, Reg1−/−, RoquinSan/san or WT mice were stimulated with TNF (10 ng/mL) and RNA expression profiles were analyzed by RT-qPCR. Data are mean ± SD.

(E) Heat map of the expression of selected LPS-inducible genes on the basis of transcriptome analysis of WT, Reg1−/−, RoquinSan/San, Reg1−/− and RoquinSan/San MEFs.

(F) GSEA of overlap between Reg1 (left) or Roquin (right) target genes (RIP-seq based) and RoquinSan/San enrichment (transcriptome analysis based). An ES plot is shown with genes ranked according to their RoquinSan/San enrichment p value. In lower panels, the ranks of Reg1 (left) or Roquin (right) target genes are indicated (blue lines).
Legends for Supplementary Tables.

Table S1. RNA sequencing data from Reg1 IP experiments, related to Figure 1

Table S2. Gene ontology (GO) annotations enriched for the 68 Reg1-interacting mRNAs, Related to Figure 1

Table S3. RNA sequencing data from Roquin-1 IP experiments, Related to Figure 2

Table S4. RNA sequencing data from Reg1 HITS-CLIP experiments, Related to Figure 2

Table S5. Ribosome is a candidate complex as Reg1-associated complex, Related to Figure 5

Proteins enriched by iTRAQ-based proteomic identifying as Reg1-associated proteins. Control or Flag-Reg1 immunoprecipitates from RAW cells was labeled with iTRAQ reagent 114 or 117, respectively. The contents of each iTRAQ reagent-labeled sample were combined and analyzed by LC-MS/MS. Based on a 80% confidence level (< 20% 117/114 variability), cutoff values of 1.2-fold for up-regulated proteins were used to define candidate proteins as Reg1-associated proteins.

Table S6. Transcriptome analysis from Reg1\(^{-/-}\); Roquin\(^{San/San}\) double mutant MEFs, Related to Figure 7

Table S7. Primers used for quantitative PCR analysis and the preparation of Northern blot probes, Related to Experimental Procedures
**A**

<table>
<thead>
<tr>
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<th>pTRE-Il6 mRNA</th>
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<td>Dox (h)</td>
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<td>Actin</td>
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**B**

Roquin (NIH3T3 cells)

**C**

Roquin (HeLa cells)

**D**

NIH3T3

**E**

Reg1 (HeLa cells)

**F**

**G**

![Graph](Graph.png)

**H**

![Graph](Graph.png)
A

Protein synthesis inhibitor

Reg1 Marker Merge

Anisomycin

Calnexin (ER)

RPL7A (Ribosome)

CHX

Calnexin (ER)

RPL7A (Ribosome)

B

Flag-Reg1

Anisomycin (10 \( \mu \)g/mL)

IB: Flag

IB: Actin

Flag-Reg1

CHX (100 \( \mu \)g/mL)

IB: Flag

IB: Actin

C

pTRE-II6 mRNA pTRE-IRES-II6 mRNA

Mock Reg1 Mock Reg1

Dox (h)

II6

Actin\( \beta \)

D

II6 mRNA

IRES-II6 mRNA

Relative remaining RNA (%)

Time (h)

Mock Reg1
A

Reg1-D141N   Marker   Merge
Calnexin (ER) RPL7A (Ribosome) TIA-1 (SG)
SA +

B

IL6 3’UTR (86-102)

Relative luciferase activity

Reg1-WT   +   +   +   Roquin-WT   +   +   +   Reg1-DN   -   -   -

Ptgs2 3’UTR-(1-2100)

Relative luciferase activity

Reg1-WT   +   +   +   Roquin-WT   +   +   +   Reg1-DN   -   -   -

C

IL6  TNF  PTGS2  NFKBIA

The ratio of RNP and Polysome RNA to total RNA (%)

D

MEFs, TNF stimulation

E

MEFs, TNF stimulation

Gene expression over time for IL6, TNF, Ptgs2, and Nfkbia under different conditions.

F

Genes enriched in Reg1 RIP-seq vs Roquin RIP-seq

Enrichment score (ES) for genes enriched in Reg1 and Roquin RIP-seq.
Table S2. Functional Categories of Reg1 target genes discovered by RIP-Sequencing analysis.

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<th>Rank</th>
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Table S5. Ribosome and UPF1 is a candidate complex as Reg1-associated complex, related to Figure 5
Proteins enriched by iTRAQ-based proteomic identifying as Reg1-associated proteins. Control or Flag-Reg1 immunoprecipitates from RAW cells was labeled with iTRAQ reagent 114 or 117, respectively. The contents of each iTRAQ reagent-labeled sample were combined and analyzed by LC-MS/MS. Based on a 80% confidence level (< 20% 117/114 variability), cutoff values of 1.2-fold for up-regulated proteins were used to define candidate proteins as Reg1-associated proteins.

**RAW cells**

<table>
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<tr>
<th>Accession No.</th>
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<th>Fold Change (117/114)</th>
<th>117/114 Variability [%]</th>
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### Table S7. Primers used for quantitative PCR analysis

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<td>Mouse Il6 (Reverse)</td>
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### Table S7. Primers used for quantitative PCR analysis (continued)

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</tr>
<tr>
<td>Human NFKBI (IkBa) (Reverse)</td>
<td>GAAGGAGATCGAGATGAGGCTC</td>
</tr>
<tr>
<td>Human GAS5 (Forward)</td>
<td>GTTCTGACTGAGCAGC</td>
</tr>
<tr>
<td>Human GAS5 (Reverse)</td>
<td>GAAGGAGATCGAGATGAGGCTC</td>
</tr>
<tr>
<td>Human SMG5 (Forward)</td>
<td>GTGACTGAGCTTACCCGCA</td>
</tr>
<tr>
<td>Human SMG5 (Reverse)</td>
<td>GTGACTGAGCTTACCCGCA</td>
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<tr>
<td>Human MAFK (Forward)</td>
<td>CGACTAATCCCCTAAGGAAGTGG</td>
</tr>
<tr>
<td>Human MAFK (Reverse)</td>
<td>GTTCTTGAGTGTGGCGGCCCA</td>
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<tr>
<td>Human STAUI (Forward)</td>
<td>CTATCTCTGAGATCGATCCAC</td>
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<td>Human STAUI (Reverse)</td>
<td>GATCCCTCAACGCTTTGTGCA</td>
</tr>
<tr>
<td>Human β-ACTIN (Forward)</td>
<td>CACAGGGAGATGGAGGCTCCTCC</td>
</tr>
<tr>
<td>Human β-ACTIN (Reverse)</td>
<td>CTCTGCTATCCTTGTGGCAATGC</td>
</tr>
<tr>
<td>Firefly Luciferase (Forward)</td>
<td>CAGAGATGGCAAGGCGCTGGA</td>
</tr>
<tr>
<td>Firefly Luciferase (Reverse)</td>
<td>CAGAGATGGCAAGGCGCTGGA</td>
</tr>
<tr>
<td>Renilla Luciferase (Forward)</td>
<td>GGATGATAACTGCTCCAGCA</td>
</tr>
<tr>
<td>Renilla Luciferase (Reverse)</td>
<td>CAGAGATGGCAAGGCGCTGGA</td>
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</table>
### List of primers (continued)

Table S7. Primers used for the preparation of Northern blot probes

<table>
<thead>
<tr>
<th>Gene</th>
<th>Sequence (5′ to 3′)</th>
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<tbody>
<tr>
<td>Mouse Il6 CDS (Forward)</td>
<td>GTTCCTCTCTGCAAGAGACTTCCATCC</td>
</tr>
<tr>
<td>Mouse Il6 CDS (Reverse)</td>
<td>GTATCTCTCTGAAGGACTCTGGCTTTG</td>
</tr>
<tr>
<td>Mouse β-Actin CDS (Forward)</td>
<td>CTATGTGGGTGACGAGCCCAGAG</td>
</tr>
<tr>
<td>Mouse β-Actin CDS (Reverse)</td>
<td>GGGTACATGGTGGTACCACCAGAC</td>
</tr>
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<td>Human UPF1 CDS (Forward)</td>
<td>CTGTGGACGACAGTGTAGCCAGAC</td>
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<tr>
<td>Human UPF1 CDS (Reverse)</td>
<td>GAGCTGTCCAGTTGATGTCCCTTGAG</td>
</tr>
<tr>
<td>Mouse β-Globin CDS (Forward)</td>
<td>CACCTGACTGATGCTGAGAAGGCTG</td>
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<tr>
<td>Mouse β-Globin CDS (Reverse)</td>
<td>CTTCTGGAAGGCAGCTGTGCAG</td>
</tr>
</tbody>
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