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Targeting TRAF6 E3 ligase activity with a small molecule inhibitor combats autoimmunity

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ABSTRACT

Constitutive NF-KB signaling represents a hallmark of chronic inflammation and autoimmune diseases. The E3 ligase TNF receptor-associated factor 6 (TRAF6) acts as a key regulator bridging innate immunity, proinflammatory cytokines, and antigen receptors to the canonical NF-κB pathway. Structural analysis and point mutations have unraveled the essential role of TRAF6 binding to the E2 conjugating enzyme Ubiquitin-conjugating enzyme E2 N (Ubc13 or UBE2N) to generate Lys63-linked ubiquitin chains for inflammatory and immune signal propagation. Genetic mutations disrupting TRAF6-Ubc13 binding have been shown to reduce TRAF6 activity and

consequently, NF-KB activation. However, to date no small molecule modulator is available to inhibit the TRAF6-Ubc13 interaction and thereby counteract NF-κB signaling and associated diseases. Here, using a highthroughput small-molecule screening approach, we discovered an inhibitor of the TRAF6-Ubc13 interaction that reduces TRAF6-Ubc13 activity both in vitro and in cells. We found that this compound, C25-140, impedes NF-KB activation in various immune and inflammatory signaling pathways also in primary human and murine Importantly, C25-140 cells. ameliorated inflammation and improved disease outcomes of autoimmune psoriasis and rheumatoid arthritis in preclinical in vivo mouse models. Hence, the

first-in-class TRAF6-Ubc13 inhibitor C25-140 expands the toolbox for studying the impact of the ubiquitin system on immune signaling and underscores the importance of TRAF6 E3 ligase activity in psoriasis and rheumatoid arthritis. We propose that inhibition of TRAF6 activity by small molecules represents a promising novel strategy for targeting autoimmune and chronic inflammatory diseases.

Ubiquitination of proteins is a cellular process that either drives protein degradation by the proteasome in order to sustain protein homeostasis or regulates distinct cellular signaling pathways. Three enzymes, namely E1, E2 and E3 are needed to covalently attach ubiquitin to the substrate proteins. E1 is the activating enzyme initially binding the ubiquitin, E2 is the conjugating enzyme thereby determining the linkage type and the E3 ligase is involved in substrate specificity and transfer of ubiquitin (1). Depending on the type of linkage, polyubiquitination can have various functions (1-3). While, e.g. Lys48-linked chains are known to target the substrate protein for degradation proteasomal via the 26S proteasome, Lys63-linked chains are associated with a variety of non-proteolytic functions such as trafficking, DNA damage response as well as NF-κB signaling in innate and adaptive immunity.

The eukaryotic transcription factor NF- κB plays a critical role in regulating the expression of a large variety of genes that are involved in several cellular processes including innate and adaptive immune response, cell growth and survival as well as cell development (4). Several signals including cytokines, pathogens, injuries and other stress conditions induce activation of the NF-kB transcription factors that are tightly regulated via the formation of poly-ubiquitin chains. Deregulation of NF-κB signaling contributes to the pathogenesis of autoimmunity, chronic inflammation and various cancers (5,6). Within NF-_KB signaling, approved ubiquitinproteasome inhibitors mainly focus on the inhibition of the 26S proteasome (e.g. Bortezomib and Carfilzomib), thereby broadly

influencing protein homeostasis. In contrast, regulators of the ubiquitination cascade might serve for more specific drug targets. Especially E3 ligases are final regulators of the ubiquitination reaction with more than 600 representatives and therefore, drugging E3 ligases might give rise to more specificity compared to the 26S proteasome (7).

The TRAF protein family comprises seven E3 ligase members including TRAF6. Different members of this class are involved in distinct receptor signaling pathways such as IL-1 receptor family, T-cell receptor (TCR), IL-17 receptor, TNF receptor and more (8,9). TRAFdependent signaling pathways contribute to the control of diverse cellular processes including differentiation survival. proliferation, and cytokine production (8). The TRAF6 protein belongs to the RING-type of E3 ligases, which interacts with the heterodimeric E2 enzyme complex Ubc13/Uev1a to attach Lys63-linked ubiquitin chains to its substrate proteins (10-12). These Lys63-linked ubiquitin chains are necessary to activate innate and adaptive immune responses mainly through the NF-kB axis (13,14). Of important note, the TRAF6 ligase activity is strictly determined by its protein-protein-interaction with a specific set of E2 enzymes including Ubc13. Importantly ubiquitin is directly transferred from ubiquitincharged Ubc13 (E2) to the substrate, both proteins bound by the E3 ligase TRAF6 (15). Thus, targeting TRAF6 E3 activity requires the of protein-protein-interaction identification inhibitors, which disrupt the TRAF6-Ubc13 interaction, because TRAF6 has no intrinsic catalytic activity (7).

Overexpression of TRAF6 as well as enhanced TRAF6 activity have been shown to promote chronic immune stimulation in a variety disorders including autoimmunity. of inflammation and cancer (8). For example, it has previously been demonstrated that TRAF6 expression is elevated in rheumatoid arthritis (RA) patients (16,17) and in Lupus cohorts (18). On a molecular level, disruption of TRAF6-Ubc13 binding by the cellular protein A20 (19) or by genetic mutations (20,21) counteracts TRAF6 activity and NF-κB activation. To date, efforts on targeting TRAF6 activity have focused on disrupting TRAF6-substrate interaction (22), whereas inhibition of the TRAF6 activity by interfering with the TRAF6-Ubc13 association has not been attempted so far and represents an attractive novel strategy to combat TRAF6-dependent disease formation.

In this report, we demonstrate the identification of the first TRAF6-Ubc13 protein-protein-interaction inhibitor that ameliorates disease outcome of two preclinical murine models of autoimmunity and chronic inflammation.

Results

Identification of TRAF6-Ubc13 inhibitors

In order to identify small molecule inhibitors of the TRAF6-Ubc13 interaction, we established an AlphaScreen[®] binding assay with robust performance (Figure S1A) and carried out a High-Throughput-Screening (HTS) campaign using 25,000 compounds (Figure S1B). After removing AlphaScreen[®] frequent hitters using our own developed chemoinformatics filters (23,24) and by employing strict selection criteria (dose-response and cell-based NF-kB activation assays), we were able to select the small molecule C25 as the most promising scaffold for evaluation (Figure further S1C-E). Subsequently, we performed a small Structure Activity Relationship (SAR) analysis by testing few commercially available analogs (Fig. S2A). Three analogs showed similar inhibitory effects on TRAF6-Ubc13 interaction when compared to C25 (Fig. S2B). Compound C25-140, however, had the best inhibitory potential on NF-KB activation after IL-1 β stimulation (Fig. S2C), which led us choose this compound for additional investigations. C25-140 (Fig. 1A and Fig. S2A) disrupted the TRAF6-Ubc13 binding in vitro (Fig. S2D), without affecting the interaction of Ubc13 and the deubiquitinase OTUB1(25) (Fig. S2D), which competes with TRAF6 for Ubc13 binding (25). Furthermore, the binding of Ubc13 to Uev1a remained unimpaired upon compound treatment (Fig. S2E). Importantly, treatment of cells with C25-140 dose-dependently impeded TRAF6-Ubc13 interaction in co-immunoprecipitation experiments (Fig. 1B) and NMR studies proved that C25-140 directly binds to TRAF6 (Fig. 1C). Furthermore, C25-140 effectively reduced TRAF6-mediated ubiquitin chain formation in *vitro* (Fig. 1D). These data demonstrate that C25-140 directly binds to TRAF6, thereby blocks the interaction of TRAF6 with Ubc13 and as a consequence lowers TRAF6 activity.

Selectivity profiling of C25-140 against other E3 ligases and E2 enzymes

To better understand the selectivity of C25-140 towards other E3 ligases, we analyzed the activity of various E3 ligases upon C25-140 treatment. These studies revealed that only the activity of cIAP1, an E3 ligase generating Lys63-linked poly-ubiquitin chains (26) similar to TRAF6, was affected by C25-140 (Fig. 2A). Notably, several other E3 ligases (MDM2, TRIM63, ITCH, E6AP and RNF4) building other types of poly-ubiquitin chains were not influenced by C25-140 (Fig. 2B-D). In addition to these studies on a set of E3 ligases, we tested whether C25-140 inhibits E1/E2 reactions. To this end, we treated different E1/E2 pairs employing nine different E2 enzymes with compound C25-140. All reactions including UbcH13/Uev1a were not affected by C25-140 (Fig. 3), confirming that C25-140 acts on the E3 ligase side.

Effects of compound C25-140 on pro-inflammatory signaling

Before moving into cell-based validation of C25-140, we analyzed whether C25-140 affects cell viability and cell cycle phases. C25-140 neither substantially influenced cell viability in MEF cells and Jurkat T-cells at concentration up to 50 μ M (Fig. S3A) nor did it influence cell cycle phases of Jurkat T-cells (Fig. S3B).

We first tested the inhibitory potential of C25-140 in MEF cells that can be stimulated with pro-inflammatory cytokines IL-1ß and TNF α to trigger NF- κ B activation. Stimulation of these cells with IL-1 β initially induces TRAF6 auto-ubiquitination that is required for signal progression through the IKK complex towards NF-KB activation and expression of NF- κ B-dependent target genes (8). C25-140 treatment of MEF cells led to a dose-dependent reduction of TRAF6 auto-ubiquitination (Fig. 4A) followed by decreased levels of phosphorylated I κ B α (Fig. 4B), which translated into diminished target gene expression (Fig. 4C).

These data illustrate that inhibition of TRAF6 activity early during IL-1 receptor signaling hampers the signaling cascade towards NF- κ B activation. Additionally, TNF α induced phosphorylation of I κ B α (Fig. S4A) as well as NF- κ B induced target gene expression (Fig. S4B) were affected by C25-140 treatment. Taken together, C25-140 efficiently inhibits IL-1 β and TNF α mediated receptor signaling in the context of cytokine activation.

Effects of C25-140 on T-cell activation

In addition to IL-1 β receptor signaling, we tested C25-140 on T-cell activation as TCR mediated NF-KB activation relies on TRAF6 activity (27,28). In the human Jurkat T-cell line, T-cell activation by PMA/Ionomycin (P/I) stimulation led to TRAF6 auto-ubiquitination that was diminished by C25-140 in a dosedependent manner (Fig. 4D). Again, decreased TRAF6 auto-ubiquitination affected downstream signaling as evident from reduced IkBa phosphorylation and cytokine secretion (Fig. 4E Moreover, P/I induced and F). JNK phosphorylation in MAPK signaling was diminished by C25-140 (Fig. S4C). All these data emphasize that C25-140 also affects TCR mediated immune response in a human derived cell line.

C25-140 is effective in cytokine signaling of primary human and murine cells

Next, we used primary human peripheral blood mononuclear cells (PBMCs) from three healthy individuals to investigate whether C25-140 is also competing TRAF6-dependent receptor signaling in primary human cells. In line with our data acquired in cell lines (Fig. 4), C25-140 reduced secretion of NF-KB-driven inflammatory cytokines such as TNF α and IL-6 after IL-1ß stimulation (Fig. 5A) or IL-1ß and TNFa upon LPS stimulation (Fig. 5B), clearly indicating that C25-140 also reduces cytokine and innate immune responses in primary human blood cells. Similar inhibitory effects by C25-140 could be proven in human PBMCs after stimulating the adaptive immune response with anti-CD3/CD28 antibodies for TCR signaling (Fig. 5C). Therefore, C25-140 is capable of inhibiting cytokine signaling also in cells of primary human origin. In addition to primary human cells, we verified the inhibitory potential of C25-140 in *ex vivo* primary murine T-cells and observed dose-dependent reduction of IL-2 mRNA and protein levels after CD3/CD28 stimulation (Fig. 5D).

Absorption/Distribution/Metabolism/Excretion (ADME) and pharmacokinetics (PK) assessment of C25-140

Before moving into *in vivo* mouse studies, we analyzed the compounds suitability for *in vivo* application. To this end, we performed ADME studies assessing key parameters (Fig. S5A). Here, C25-140 was stable in plasma as well as in microsomes, showed 96.8% plasma protein binding and exhibited low binding to hERG. Only in terms of Cytochrome P450, the compound showed some inhibitory potential on distinct CYPs (Fig. S5A).

Next, we tested the pharmacokinetics (PK) of C25-140 after intravenous (IV), peroral (PO) and intraperitoneal (IP) injection of mice with 10 mg/kg C25-140 each. In general, a rapid initial distribution phase was observed and C25-140 had good calculated oral bioavailability (Fig. S5B and C).

Altogether, results from ADME and pharmacokinetics measurements underscored that compound C25-140 had appropriate properties to be applied in mouse studies for *in vivo* efficacy testing.

C25-140 efficacy in a psoriasis mouse model

In a first *in vivo* mouse model we aimed at investigating the effects of C25-140 on psoriasis, an autoimmune disease. In this model, psoriasis is induced by imiguimod (IMO), which activates TLR7 signaling (29), a pathway employing TRAF6 for signal progression. IMQ was topically applied to the shaved back as well as the right ear of mice. On the same regions C25-140 was topically applied twice daily at a final dose of approx. 1.5 mg/kg per application. Parameters describing disease outcome were scored every day and samples for IL-17 cytokine measurement were collected at day 6 (Fig. 6A). C25-140 was able to significantly reduce the "cumulative score" (Figure 6B), the overall "thickness score" (Fig. 6C), "scaling" (Figure 6D) and "erythema" (Fig. 6E). Moreover,

analysis of IL-17 cytokine expression in the right ear tissue proved its downregulation as a consequence of C25-140 treatment (Fig. 6F). This first efficacy study demonstrates that C25-140, an inhibitor of TRAF6 activity, is able to ameliorate symptoms of autoimmune psoriasis.

C25-140 efficacy in a preclinical mouse model for rheumatoid arthritis (RA)

As pointed out earlier, it has previously been reported that levels of TRAF6 are elevated RA patients (16,17). Therefore, we in investigated whether C25-140 is effective in a preclinical mouse model of autoimmune rheumatoid arthritis. We chose the collageninduced arthritis (CIA) mouse model (30) in which TRAF6 is upregulated during disease progression and RA symptoms are reversed by the application of siRNA-based TRAF6 knockdown (17), stressing the biological relevance of this in vivo model for the TRAF6 E3 ligase functions. In our study, RA was induced by injection of collagen at day 0, followed by a collagen boost injection at day 21 to promote the development of arthritic symptoms. At day 28, compound C25-140 was intraperitoneally administered twice daily at three doses (6mg/kg, 10mg/kg and 14mg/kg) over a period of 14 days. In RA, patients usually receive treatment when symptoms become apparent. Thus, we started dosing C25-140 in the CIA model when mice already had developed symptoms of RA (arthritic index of approx. 3), thereby reflecting the diseased patient. Prednisolone served as a positive control. Mice were daily scored for arthritic index (AI) and euthanized on day 42 in order to collect limbs for histopathology (Fig. 7A). Throughout the entire study the body weight of all mice was monitored and did not show any reduction, indicating that there were no signs of obvious toxicity (Fig. S6). Intriguingly, C25-140 ameliorated the arthritic index to almost baseline levels in this efficacy model at the doses of 10mg/kg and 14mg/kg (Fig. 7B). Tissue sections of the limbs (Fig. 7C and Fig. S7) evidently verified that C25-140 dose-dependently improved symptoms of RA including inflammation and structural damages. In addition, all sections were quantified based on parameters of disease appearance such as summed scores, inflammation, pannus, cartilage

damage and several more (Fig. 7D and E and Fig. S8). These data also underlined the dosedependent improvement of RA disease outcome. Taken together, we present a successful preclinical proof of concept study for C25-140 in a RA efficacy model.

Discussion

In this report, we present a highthroughput screening campaign that successfully discovered a novel protein-protein-interaction inhibitor, namely C25-140, which is capable of reducing TRAF6 E3 ligase activity by interfering with the TRAF6-Ubc13 interaction. Moreover, C25-140 exhibits inhibitory effects on various immune and inflammatory signaling pathways in cell lines as well as primary human and murine cells. Notably, inhibition of these pathways should be beneficial for treating a highly inflamed immune system. In line with this notion, C25-140 ameliorated symptoms of autoimmune diseases in in vivo preclinical efficacy models of psoriasis and rheumatoid arthritis (RA). Hence, our reverse chemical genetics approach presented in this study underlines the importance of TRAF6 activity for the development and persistence of autoimmune diseases such as psoriasis and RA.

TRAF6 belongs to the RING type of E3 ligases, where the RING domain makes a protein-protein-interaction with the ubiquitincharged Ubc13 protein. Here, the ubiquitin is transferred from Ubc13 (E2) to the substrate and TRAF6 acts as a scaffold bridging these two proteins (15). Therefore, targeting the E3 activity of TRAF6 requires protein-proteininteraction inhibitors, which interfere with the binding of TRAF6 to Ubc13 (7). Proteinprotein-interaction inhibitors (PPIs) have been the very difficult to target in drug discovery as the interaction interfaces are usually large and contain many hydrophobic residues (31-33). However, PPIs comprise hotspot regions and genetic point mutation can lead to complete disruption of binding, thereby suggesting that small molecules have the chance to interfere with these interfaces (33). Indeed, crystal structures of TRAF6-Ubc13 binding clearly imply that distinct amino acids of TRAF6 (e.g. C70 or D57) are crucial for its interaction with Ubc13 (10). Single point mutations of these

amino acids entirely abrogated TRAF6-Ubc13 interaction (Figure S1A and (10)), emphasizing that there is a good chance to impede the TRAF6-Ubc13 PPI.

The goal of this study was to identify a selective inhibitor of TRAF6 activity that can be applied to chemical biology as well as drug discovery studies. Our selectivity profiling on other E3 ligases including RING (e.g. cIAP1 and MDM2) and HECT (e.g. ITCH and E6AP) type of E3 ligases revealed that C25-140 comprises good selectivity for TRAF6 over other E3 ligases with the exception that it also inhibits cIAP1, a RING E3 ligase generating Lys63 ubiquitin chains, similar to TRAF6. Importantly, C25-140 did not interfere with Lys48-linked poly-ubiquitin chain formation, which is critical to maintain cellular protein homeostasis. The compound also did not inhibit several other E1/E2 reactions including UbcH13, underlining that it acts on the E3 ligase side of the PPI. Hence, C25-140 is fairly selective and does not act as a broadband inhibitor of E3 ligases.

The inhibition of TNF α signaling by C25-140 was unexpected, as this particular receptor activation does not employ TRAF6 for signal progression towards NF- κ B (34). This finding, however, can be explained by the fact that C25-140 also inhibits cIAP1 E3 ligase activity, which is a central regulator of TNFa receptor signaling (26). In fact, this C25-140 offtarget activity may eventually be beneficial for the treatment of autoimmune and inflammatory diseases as uncontrolled function of $TNF\alpha$ signaling has been shown to highly contribute to the development of these diseases (35). Therefore, our study provides a novel small counteracts TRAF6-Ubc13 molecule that interaction as a target to reduce NF-kB signaling by various receptors (IL-1ß receptor, TLR4, Tcell receptor and TNF α receptor), thereby encompassing cumulative effects on signaling processes that drive chronic inflammation.

Current therapeutics for RA are mostly "disease modifying anti-rheumatic drugs" (DMARDs) and glucocorticoids with broad and diffuse immunosuppressive effects in the early stage (36-38) with addition of antibody/biologics therapy against specific surface targets (39,40) at later stages. The only approved small molecule with target-based function against RA is Tofacitinib, a JAK inhibitor (41-43), which yet comprises serious side effects. Thus, the field relies on the discovery of additional approaches that use small molecules against novel diseaserelated targets. Accordingly, C25-140 is a novel small molecule inhibitor that interferes with a distinct target, i.e. TRAF6 activity, which is critical for activation of various NF- κ B signaling pathways. This novel strategy effectively leads to reduction of pro-inflammatory signaling to improve RA disease outcome.

In the past, some inhibitors of MYD88mediated receptor signaling have been described including IRAK4 activity (44), Ubc13-Uev1a interaction (45), MYD88 dimerization (46) and Ubc13 (47), but targeting the E3 ligase activity of TRAF6 has never been accomplished as its ligase activity is determined by the E2-E3 protein-protein-interaction. In general, inhibition of E2-E3 interactions have been challenging and only inhibitors of an acetylated-E2 together with a Cullin type E3 ligase have been reported (48). In contrast, our study describes the first-in-class protein-protein-interaction inhibitor of a RING-E3 ligase (TRAF6) / E2 enzyme (Ubc13) binding and presents a novel strategy to treat autoimmune and chronic inflammatory diseases.

Experimental procedure

Plasmids, cell lines, antibodies and compounds Cell lines:

HEK293T and MEF were grown in DMEM (Gibco) supplemented with 10% Fetal bovine serum (FBS) (Gibco) and 100 U/ml Penicillin/ Streptomycin (Gibco). Jurkat cells were grown in RPMI 1640 (Gibco) supplemented with 10% FBS (Gibco) and 100 U/ml Penicillin/Streptomycin (Gibco). Cells were rethawed every three to four weeks. Human PBMCs were cultured in RPMI supplemented with 10% FBS, 100U/ml Penicillin/ Streptomycin and 50 µM b-Mercaptoethanol (Gibco). Primary mouse CD4+ T-cells were cultured as previously described (49).

Plasmids:

cDNA of TRAF6 RZ 12 (residues 50-187) was cloned into pEF-HA backbone vector (50). cDNA of TRAF6 RZ1 (residues 50-159) or the TRAF6 D57K mutant was cloned into pASK IBA 3+ (IBA Lifesciences) and pGEX4T1 (GE Healthcare). Full length Ubc13 was cloned into the pGEX4T1 vector with and without Cterminal Flag-His-tag. OTUB1 full-length cDNA was cloned into pGEX4T1 and Uev1a was cloned into pET28a vector.

Antibodies:

These antibodies were used for Western Blotting: β -Actin I-19 (Santa Cruz sc-1616, 1/1000); p-I κ B α Ser32/36 5A5 (Cell signaling #9246, 1/1000); I κ B α C21 (Santa Cruz sc-371, 1/1000) for MEF cells; I κ B α L35A5 (Cell Signaling #4814, 1/1000) for Jurkat T-cells; RNF4 (Abnova A01, 1/2000); TRAF6 (Abcam EP591Y, 1/1000); Ubc13 (Invitrogen 37-1100, 1/1000); Ubiquitin P4D1 (Santa Cruz sc-8017, 1:1000).

Anti-HA Affinity Matrix (Roche Diagnostics, 1/40) was used for immunoprecipitation.

Compounds:

All compounds were purchased from ChemDiv: C25 (Cat. No.: G827-0171) C25-140 (Cat. No.: G827-0140) C25-167 (Cat. No.: G827-0167) C25-189 (Cat. No.: G827-0189) C25-031 (Cat. No.: G827-0031) C25-166 (Cat. No.: G827-0166)

C25-054 (Cat. No.: G827-0054)

C25-024 (Cat. No.: G827-0024)

For all experiments, C25-140 was re-synthesized to ensure quality control for working with the correct molecule (synthesis procedure below).

Recombinant protein expression and purification

Purification of Strep-tagged TRAF6 RZ1 protein The plasmid was transformed into BL21 codon plus RIPL cells (Agilent Technologies) and cultured in Luria/Miller medium containing Ampicillin and Chloramphenicol. Protein production was induced at an OD600 = 0.6-0.8 by adding 200 µg anhydrotetracycline, 0.5 mM Isopropyl-β-D-thiogalactopyranosid (IPTG) and 100 µM ZnCl₂. Cells were incubated at 18°C for 16 hours. Afterwards, cells were centrifuged at 2,600 xg for 20 minutes at 4°C. The pellet was resuspended in 10 ml Strep-wash buffer (IBA Lifesciences) including 1 pill of protease

inhibitor Complete mini (Roche Diagnostics). The suspension was sonified (8 x 30 seconds) and subsequently centrifuged at 53,000 xg for 2 x 30 minutes. For the purification of the StrepIItagged proteins, the protein solution was applied to Strep-Tactin columns using the ÄKTA purifier (GE Healthcare). After washing, the StrepII-protein was eluted from the column using the Strep-elution buffer (IBA Lifesciences) containing desthiobiotin. The eluted protein was concentrated using Amicons (cut off 3 kDa) down to 1 ml. Using the ÄKTA purifier and a HiTrap Desalting 5 ml column (GE Healthcare), the protein was desalted and exchanged to storage buffer (20 mM Tris-HCL pH 8.0; 20 mM NaCl, 1 mM 1,4 Dithiothreitol and 100 µM ZnCl₂). The protein was concentrated in Amicons up to 2 µg/µl and stored in small aliquots at -80°C.

Purification of untagged TRAF6 RZ1 protein

The cDNA sequence of wildtype TRAF6 RZ1 domain was cloned into the pGEX4T1 vector (GE Healthcare Life Sciences). For the purification of untagged TRAF6 RZ1, the protein was expressed in BL21 codon plus RIPL cells in M9 minimal medium. For ¹⁵N-labeled protein, the regular NH₄ was replaced by ¹⁵NH₄. For protein production, 1 mM IPTG and 100 µM $ZnCl_2$ was added at OD600 = 0.6-0.7. After 16 hours at 18°C, cells were centrifuged and the pellet resuspended in 10 ml TRAF6 washbuffer (20 mM Tris pH 8.0, 20 mM NaCl) including 1 Complete mini pill (Roche Diagnostics). After sonification (8 x 30 seconds) and centrifugation (2 x 30 minutes at 53,000 xg), the protein suspension was incubated with 1 ml pre-washed Glutathione Sepharose 4 fast flow beads (GE Healthcare) for 2 hours at 4°C. Subsequently, beads were washed with 100 ml TRAF6 wash buffer and then incubated with 50 Units Thrombin for 3 hours at room temperature to cleave off the GST-tag. The TRAF6 protein was eluted in 5 ml wash buffer and cleared for remaining GST with 1 ml Glutathione Sepharose 4 fast flow beads. Afterwards, the solution was centrifugated for 5 minutes at 200 xg and the supernatant transferred into an Amicon for concentration followed by centrifugation at 20,000 xg for 30minutes. The supernatant was then applied to a HiLoad Superdex75 column

(GE Healthcare) using the ÄKTA purifier for separating the untagged TRAF6 protein from uncleaved protein as well as buffer exchange in desalting buffer (2 mM Tris pH 8.0, 20 mM NaCl, 1 mM DTT and 100 μ M ZnCl₂). These proteins were used for 2D-NMR and *in vitro* ubiquitination experiments.

Flag-His-tagged Ubc13 protein (Ubc13-FH) and GST proteins (including GST-OTUB1 and GST-Ubc13) were produced as previously described (51).

His-tagged Uev1a protein was purified according to GST-OTUB1 protocol with the following buffer: lysis buffer (20 mM NaH₂PO₄, 20 mM NaCl, 50 mM Imidazol), elution buffer (20 mM NaH₂PO₄, 20 mM NaCl, 500 mM Imidazol) and desalting buffer (20 mM NaH₂PO₄, 20 mM NaCl). Ni-Sepharose 6 Fast Flow (GE Healthcare) was used instead.

AlphaScreen[®] assay for compound screening of TRAF6-Ubc13 PPI inhibitors

For performance of an automated AlphaScreen[®] assay, all following components were prediluted in AlphaScreen® buffer (1x PBS, 0.5 % BSA, 0.01 % Tween-20). First, the TRAF6 protein (end concentration 100 nM, 30 µl volume) was added to 384-well Opti-plates (PerkinElmer) involving the MultiFlo dispensing system (BioTek) followed by transfer of the compounds via the Sciclone G3 transfer station (PerkinElmer). After addition of Ubc13FH (end concentration 75 nM, 10 µl volume) with the MultiFlo system and incubation for one hour at room temperature, both beads (Strep-Tactin Alpha Donor beads and Nickel-Chelate acceptor beads; end concentrations 4 µg each, 10 µl volume each, PerkinElmer) were added using the MultiFlo system in subdued light. Read-out of the plates occurred after another hour of incubation at room temperature. Statistical parameters including the coefficient of variation (%CV), Z' factor and signal window (SW) were calculated to determine the quality of the assay. To evaluate the efficacy of the compounds, AlphaScreen® units of compound-treated samples were referred to DMSO control treated samples. Thereby, the TRAF6_{D57K} mutant served as the control for minimum signal and was

included on every plate of the screening. Compounds that inhibited TRAF6_{WT}StrepII-Ubc13FH by more than 25 % were considered as actives. After elimination of AlphaScreen[®] frequent hitter compounds and His-tag frequent hitters(23), 178 compounds were defined as primary hits and were subsequently tested in five-point serial dilution assays $(2.5 - 40 \mu M)$ in TRAF6_{WT}StrepII–Ubc13FH AlphaScreen experiments. Only compounds with doseeffects on TRAF6_{WT}StrepIIdependent Ubc13FH (n = 27) were further taken.

AlphaScreen® counter assays_involved the interactions of GST-OTUB1/Ubc13FH and GST-Ubc13/Uev1a-His. For each assay, the GST-tagged protein was applied at a final concentration of 30 nM (30 μ l), wherein the Histagged protein was used at a concentration of 20 nM (10 μ l). Glutathione-Donor beads and Nickel-Chelate Acceptor beads (Perkin Elmer) were used at a final concentration of 3 μ g/ml (10 μ l) each).

Two-dimensional-Nuclear-Magnetic-Resonance (2D-NMR)

To determine whether the inhibitory compounds directly bind to TRAF6 rather than to Ubc13. experiments 2D-NMR were conducted. Untagged ¹⁵N-labeled TRAF6 RZ1 protein (c =120 μ M) and compound (c = 20 mM stock) were incubated at a ratio of 1:5 in a 3 mm NMR tube for 10 minutes before spectra acquisition. To observe chemical shift perturbations upon compound addition, two-dimensional SOFAST-HMOC spectra were acquired. All measurements were performed using the Bruker Avance 600 MHz spectrometer.

In vitro Ubiquitination assays (Western Blotbased)

The ability of TRAF6 to form K63-linked polyubiquitin chains in conjunction with Ubc13 was analyzed after compound treatment. Therefore, untagged TRAF6_{WT} as well as TRAF6_{D57K} were recombinantly purified. 0.125 μ M of TRAF6 protein was pre-incubated with DMSO or compound in a total volume of 100 μ l in K63 assay buffer (25 nM Tris-HCl pH 7.6; 250 nM MgCL₂; 500 nM creatine phosphate; 0.3 U/mL inorganic pyrophosphatase; 0.3 U/mL creatine phosphokinase) for 30 minutes at room temperature. Small aliquots for input samples were taken. A master mix containing 0.01 µM E1-activating enzyme (UBE1) (Boston Biochem), 0.2 µM E2-conjugating enzyme complex (Ubc13/Uev1a) (Boston Biochem), 1 mM ZnCl₂, 2 mM ATP and 4 µM monoubiquitin (Boston Biochem) was added to the TRAF6 protein and the reaction mixture was incubated for 120 minutes at 37°C. The reaction was stopped by adding 4x SDS loading buffer and boiling at 95°C for 5 minutes. The input samples were analyzed using the Pierce Silver (Thermo Stain Fisher Scientific). Kit Ubiquitination was detected in Western Blot analysis using an ubiquitin antibody (52).

For analysis of cIAP1, MDM2, TRIM63, RNF4, ITCH and E6AP mediated poly-ubiquitination, the respective ubiquitin ligase kit from Boston Biochem (K-102, K-260, K-200B, K-220, K-240 and K-270) was used. Here, reaction buffer, E1 activating enzyme, E2 conjugating enzyme and E3 ligase and substrate (p53 for MDM2, S5A for TRIM63 and S5A for E6AP) were mixed and pre-incubated with compound or DMSO for 5 minutes at room temperature before adding Mg²⁺-ATP and mono-ubiquitin for reaction start. Ubiquitination was visualized by Western Blot as described before (52).

E1/E2 reactions

UbcH (E2) enzyme kit (K-980B) and E1 enzyme (UBE1, e-305) were purchased from Boston Biochem. Ubiquitination assays were set as described above for cIAP1 but only involving E1/E2 enzymes. After 30 min of incubation at 37°C, the reaction was stopped and analyzed in Western Blot staining for Ubiquitin P4D1.

Cell culture methods

AlphaSurefire

In order to detect protein levels of p-I κ B α (Ser32/36) as well as total I κ B α , 1x10⁴ MEF cells were seeded in 96-well plates, treated with compound for 6 hours and stimulated with 1 ng/ml IL-1 β for 7 minutes. The p-I κ B α and total I κ B α protein levels were analyzed using the Alpha*Surefire* kits (PerkinElmer) following the manufacturer's instruction. The general 2-plate protocol was conducted.

Co-Immunoprecipitation studies in HEK 293T cells

To study the interaction of TRAF6 and Ubc13 in cells, 1×10^6 293T cells were seeded and after 18 hours cells were transfected with 1 ug pEF-HA empty or pEF-HA-TRAF6 RZ1 plasmid using the X-tremeGENE HP DNA transfection reagent (Roche Diagnostics). 6 hours upon transfection, cells were treated with compound or DMSO. After 42 hours of further incubation, cells were harvested in Co-IP lysis buffer (150 mM NaCl; 25 mM Hepes (pH 7.5); 0.2% NP-40; 1 mM Glycerol; 10 mM NaF; 8 mM β-Glycerophosphat; 1 mM DTT; 300 µM Sodiumvanadate: Complete protease inhibitor) and HA-TRAF6 was immunoprecipitated with anti-HA Affinity Matrix (Roche Diagnostics). Binding of Ubc13 to TRAF6 was examined by Western Blot.

Immunoprecipitation of TRAF6 for detection of endogenous ubiquitination

 2.5×10^6 MEF cells were seeded, treated with compound for 6 hours and stimulated with IL-1B (3.5 ng/ml) for 10 minutes. For Jurkats T-cells, 2.5×10^6 cells were seeded, treated with compound for 6 hours and stimulated with PMA (400 ng/ml) and Ionomycin (600 ng/ml) (P/I) for 10 minutes. Cells were lysed in Co-IP lysis buffer containing 1 % SDS to eliminate all cellular interactions. After lysis, centrifugation and collection of input samples, the samples were diluted to 0.1 % SDS and incubated with 5 ul TRAF6 antibody EP591Y (Abcam) and immunoprecipitated with Protein-G-Sepharose beads (GE Healthcare). After washing of beads in Co-IP wash buffer (150 mM NaCl; 25 mM Hepes (pH 7.5); 0.2% NP-40; 1 mM Glycerol), TRAF6 protein was eluted and levels of TRAF6 ubiquitination were detected by Western Blot.

Densitometric quantification of Western Blot bands

Western blots were densitometricaly quantified using the LabImage1D software provided by Kapelan Bio-Imaging. Here, backgroundsubtracted values of the protein of interest were calculated relative to the background-subtracted values of the control protein.

Analysis of mRNA expression

For MEF cells, 1.2×10^5 cells per sample were seeded, treated with compound for 6 hours and stimulated with either IL-1 β (1 ng/ml) or TNF α (10 ng/ml) for 60 minutes. Jurkat T-cells and primary mouse CD4⁺ T-cells were seeded at 2 x 10^5 cells per sample, also treated with compound for 6 hours and stimulated with P/I for 3 hours. RNA was isolated according to the manufacturer's protocol of the Qiagen RNeasy Kit (MEF cells) or the STRATEC INVITRAP® SPIN UNIVERSAL RNA MINI KIT (Jurkat Tcells). To clear isolated RNA from genomic DNA, samples (up to 5µg RNA) were treated with RO1 RNase-Free DNaseI (Promega) as described in the technical manual. For the reverse transcription of RNA into cDNA, the DNaseI digested RNA samples were processed using the SuperScript III First Strand cDNA Synthesis System (Invitrogen/ThermoFisher Scientific) the following manufacturer's protocol. Here, random hexamers provided with the kit were used for reverse transcription. cDNA samples were stored at -20°C. For quantification of cDNA in the samples, the LightCylcer480 (Roche) and the KAPA SYBR FAST qPCR Kit were used as described earlier(52). For relative quantification, the $\Delta\Delta Cp$ method first described by Pfaffl (53) was used. The following primer were used for qRT- PCR:

Mouse A20 for	5' GCT CAA CTG GTG TCG
	TGA AG 3'(SEQ ID NO:5)
Mouse A20 rev	5' ATG AGG CAG TTT CCA
	TCA CC 3'(SEQ ID NO:6)
Mouse β-Actin	5' CCT CTA TGC CAA CAC
for	AGT GC 3' (SEQ ID NO:7)
Mouse β -Actin	5' GTA CTC CTG CTT GCT
rev	GAT CC 3' (SEQ ID NO:8)
Mouse ICAM-1	5' CGC TCA GAA GAA CCA
for	CCT TC 3'(SEQ ID NO:9)
Mouse ICAM-1	5' GGA GAC GCA GAG GAC
rev	CTT AAC 3'(SEQ ID NO:10)
Mouse IL-2 for	5' GAG TGC CAA TTC GAT
	GAT GAG 3' (SEQ ID NO:13)
Mouse IL-2 rev	5' AGG GCT TGT TGA GAT
	GAT GC 3' (SEQ ID NO:14)
Human IL-2 for	5' CAC AGC TAC AAC TGG
	AGC ATT TAC 3'(SEQ ID
	NO:17)

Human IL-2 rev	5' TGC TGA TTA AGT CCC
	TGG GTC 3'(SEQ ID NO:18)
Human TNFα	5' CCC AGG GAC CTC TCT
for	CTA ATC 3'(SEQ ID NO:19)
Human TNFα	5' GCT ACA GGC TTG TCA
rev	CTC GG 3'(SEQ ID NO:20)
Human RNA	5' GCA CCA CGT CCA ATG
Pol II for	ACA 3'(SEQ ID NO:21)
Human RNA	5' GTG CGG CTG CTT CCA
Pol II rev	TAA 3'(SEQ ID NO:22)

Enzyme-Linked ImmunoSorbent Assay (ELISA) For measurement of cytokine secretion, 2×10^5 Jurkat T-cells or primary mouse CD4⁺ T-cells or human PBMCs were seeded per sample, subsequently treated with compound for 6 hours and stimulated as indicated. 20 hours after stimulation, supernatants were harvested and analyzed with the IL-2, IL-1 β , IL-6 and TNF α Ready-Set-Go! ELISA (2nd Generation) kits provided by Affymatrix eBioscience.

Viability, cell cycle, ADME, PK and safety studies

Viability assay

For cell viability experiments the CellTiter-Glo® 2.0 system (Promega) was used. MEF cells (1.2×10^3 cells/well) and Jurkat T-cells (5×10^3 cells/well) were seeded in 384 well plates and 18 hours later, cells were treated with compounds using the Sciclone G3 automation system. After 24 hours incubation, the luciferase assay was performed according to the manufacturer's manual. Signals were measured on an Envision plate reader (PerkinElmer).

Cell cycle analyses

For cell cycle staining, cells were seeded at 2 x 10^5 cells per sample and treated with compound or DMSO. After the indicated incubation time (24 or 72 hours), cells were fixed in 100 % Ethanol and subsequently stained with the Propidium Iodide staining solution (0.1 % Triton X-100; 20 µg/ml Propidium Iodide; 200 µg/ml RNAse A in PBS) for 40 min in the dark. The Propidium Iodide fluorescence was collected using the Attune Flow Cytometer (Thermo Fisher Scientific) and analysis of cell cycle phases was performed using the Single Cell analysis software FlowJo as previously published (54).

ADME and pharmacokinetic studies

The Bienta Enamine Biology Services of the CRO Bienta/Enamine Ltd. conducted all ADME (plasma stability, plasma protein binding, microsomal stability, LogD, Caco-2 assay, CYP450 inhibition and hERG predictor assay) as well as pharmacokinetic studies (T_{max}, C_{max}, AUC_{0-240min}, Mean Residence Time, Elimination half life, Elimination rate constant, Volume of Distribution and Clearance) of C25-140 by using their standard protocols. Pharmacokinetic measurements were done by intraperitoneal (IP), intravenous (IV) or peroral (PO) application of 10 mg/kg C25-140. Each time point was investigated with n = 4 mice. Bioethical permission of pharmacokinetic measurements in mice was granted under the numbers: IACUC number №1 from 19.05.2015 and IACUC number No6 from 08.12.2015 to Bienta.

Mouse studies

Isolation of primary mouse CD4 positive T-cells (ex vivo)

 $CD4^+$ T-cells from the peripheral lymph nodes of Balb/C mice were isolated by negative magnetic-activation cell sorting (MACS) selection using the $CD4^+$ T-cell isolation kit II (Miltenyi). Per sample, 2 x 10⁵ cells were seeded and treated with compound for 6 hours. Cells were stimulated with anti-CD3 (0.5 mg/ml) and anti-CD28 (1 mg/ml) on plates that were precoated with rabbit anti-hamster IgG. Samples for qRT-PCR were harvested after 4 hours whereas supernatants for analysis of cytokine secretion were harvested 20 hours after stimulation.

Psoriasis mouse study (in vivo)

To evaluate the anti-inflammatory activity of the compound C25-140, an Imiquimod (IMQ)induced psoriasis mouse model was carried out by the CRO Washington Biotechnology, Inc. (Baltimore, USA). Per group (vehicle and C25-140 treated), 8 male Balb/C mice were shaved in the back (1.5 cm x 2 cm) and IMQ and C25-140 were applied topically to the shaved back and the right ear every day for 6 days. Here, C25-140 was dissolved in acetone and applied at a total dosage of 500 μ g per day (2 x 250 μ g). This refers to a total of approx. 1.5 mg/kg per application. Mice were monitored and daily scores were independently collected on a scale from 0 to 4: 0=none; 1=slight; 2=moderate; 3=marked and 4=very marked. Ear thickness was measured by electronic calipers as an indicator of edema. IL-17 cytokine secretion of the right ear tissue was measured by ELISA. Bioethical permission of the IMQ-induced psoriasis study was granted under the IACUC number 17-063 to WB.

Collagen-Induced Arthritis Model (in vivo)

In order to test the potential of C25-140 to ameliorate RA symptoms, a Collagen-Induced Arthritis (CIA) Model in DBA1/J mice was performed by the CRO Washington Biotechnology, Inc. (Baltimore, USA). Each group of 10 mice was subjected to a single subcutaneous injection of collagen/complete Freund's Adjuvant emulsion (0.05 ml/mouse; 100 µg collagen/CFA). After 20 days (day 21), mice were booster injected with collagen (0.05 ml/mouse; 100 µg/mouse collagen in Incomplete Freund's Adjuvant). On day 28, mice were scored for their Arthritic Index (AI) and 14 days intraperitoneally dosed twice daily with vehicle; 6 mg/kg C25-140, 10 mg/kg C25-140 and 14 mg/kg C25-140. 10 mg/kg prednisolone was applied once daily. Mice were daily scored for macroscopic signs of arthritis. Thereby, the following scoring index was applied to each individual paw: 0 = no visible effects of arthritis; 1 = edema and/or erythema of 1 digit; 2 = edema and/or erythema of 2 digits; 3 = edema and/orerythema of more than 2 digits; 4 = severe arthritis of entire paw and digits. Single paw scores were added and recorded. On day 42, animals were euthanized, limbs were collected and histopathology was performed using Toluidine blue staining with scoring (0-5) of several parameters (summed score. inflammation, pannus, bone resorption, cartilage damage, periosteal bone formation and bone width). Bioethical permission of the collageninduced arthritis study was granted under the IACUC number 17-059 to WB.

Experiments with human primary PBMCs

For isolation of peripheral blood mononuclear cells (PBMCs) from 50 ml human blood of three different donors, blood was mixed with 800 units of Heparin (Sigma-Aldrich) and centrifuged at 300 xg for 10 minutes without brake. After removal of the plasma fraction, buffy coat was diluted in 2 volumes PBS and added on top of 15 ml Lymphoprep (STEMCELL technologies). After centrifugation at 160 xg for 20 minutes at room temperature without brake, platelets were removed and the cell suspension was centrifuged again at 350 xg for 20 minutes without brake. Mononuclear cells from the intermediate layer were transferred to a new falcon and washed 3 times in PBS supplemented with 0.1 % BSA and 2 mM EDTA. In the end, cells were resuspended in RPMI medium containing 10% FCS, 1% Penicillin/Streptomycin and 50 μM βmercaptoethanol. Per sample, 2×10^5 cells were seeded and treated with compound for 6 hours. The cells were stimulated with either 1 µg/ml Lipopolysaccharide (LPS) (Sigma-Aldrich); 20 ng/ml IL-1b (R&D Systems) or CD3/CD28 (1 μ g hCD3 (IgG_{2A}), 4 μ g hCD28 (IgG₁), 2 μ g ant-IgG_{2A} and 2 μ g ant-IgG_{2A}; all BD Pharmingen). Supernatants for analysis of cytokine secretion were harvested 20 hours after stimulation. The permission to isolate, treat and analyze human peripheral white blood cells ex vivo was granted by the ethics commission of the Technical University Munich (TUM) (project number 358/15).

Ethical statement Mouse studies:

- Bioethical permission of pharmacokinetic measurements in mice was granted to Bienta/Enamine under the numbers: IACUC number №1 from 19.05.2015 and IACUC number №6 from 08.12.2015.
- Bioethical permission of the IMQ-induced psoriasis study was granted to Washington Biotechnology, Inc. under the IACUC number 17-063.
- Bioethical permission of the CIA-induced RA study was granted to Washington Biotechechnology, Inc. under the IACUC number 17-059.

Human ex vivo studies (PBMCs):

The permission to isolate, treat and analyze peripheral white blood cells from healthy donors *ex vivo* was granted by the **ethics commission** of the **Technical University Munich (TUM)** (project number 358/15). The studies abide by the Declaration of Helsinki principles.

Synthesis procedure C25-140 General Methods:

All solvents were purchased from Sigma-Aldrich and were used as received; anhydrous solvents were used for reactions, and HPLC grade solvents were used for aqueous recrystallizations work ups. and chromatography. Other reagents were purchased from various vendors and were used as received. Glassware was dried by keeping it in an oven at 70 °C for 12 h prior to use. The pH of aqueous solutions was estimated using pH paper. Vacuum filtrations were carried out using a house vacuum line. In the individual procedures, the phrase "concentrated in vacuo" means that solvent was removed on a rotary evaporator using a diaphragm pump (with an automatic vacuum regulator) and then remaining traces of volatiles were removed on a high-vacuum (<1 torr) oil pump.

Reactions were monitored by TLC using EMD silica gel 60 F254 (250 µm) glass-backed plates (visualized by UV fluorescence) and by LC-MS Waters Acquity H UPLC CLASS system, with Aquity UPLC BEH C18 column (1.7 µm, 2.1x50 mm), eluting at 0.8 ml/min, using a 3 min linear gradient method with a mobile phase consisting of water/acetonitrile $(0.05 \% \text{ v/v TFA added to each}): 95:5 \rightarrow 5:95 (0-$ 2.25 min), 95:5 (2.27-3 min). Sample runs were monitored using alternating positive/negative electrospray ionization (50-1000 amu) and UV detection at 254 nm. Automated preparative normal-phase chromatography was carried out on a Büchi Reveleris Prep. Purification system with 3 channel viable UV-Vis and ELS detection (runs were monitored at 220-400 nm) Prepacked silica gel cartridges (12, 25 and 40 g) employed normal-phase were for chromatography, eluting at 20-30 ml/min. ¹H NMR spectra were recorded at 400 MHz on a Bruker spectrometer and are reported in ppm using the residual solvent signal (dimethylsulfoxide- $d_6 = 2.50$ ppm) as an internal standard. Data are reported as: $\{(\delta \text{ shift}),$ [(s=singlet, d=doublet, dd=doublet of doublets, br=broad, m=multiplet), (J=coupling constant in Hz) and (integration)]}.



1-(3-Methyl-[1,2,4]-triazolo-[4,3-b]-pyridazin-6-yl)-hydrazine. A round-bottomed flask was charged with 6-chloro-3-methyl-[1,2,4]-triazolo-[4,3-b]-pyridazine (2.5 g, 0.015 mol) and ethanol (35 mL). To the resulting solution was added the hydrazine hydrate (2.6 mL, 0.053 mol). The reaction mixture was then heated at 80°C for 3h. After cooling to room temperature, the formed product slurry was filtered, washed with little ethanol and air dried to afford 2.43g (98%) of the desired product as a white solid. ¹H NMR (400MHz, DMSO-d₆): δ 8.46 (br s, 1H), 7.86 (d, J = 10 Hz; 1H), 6.78 (d, J = 10 Hz; 1H), 4.28 (br s, 2H), 2.55 (s, 3H); LC-MS (ESI+) *m/z*: $[M+H]^+$ Calcd. for C₆H₉N₆ 165.17 found 165.20, $t_{\rm R} = 0.17$ min.



3-(3,5-dimethyl-1-(3-methyl-[1,2,4]-Methvl triazolo-[4,3-b]-pyridazin-6-yl)-1H-pyrazol-4yl)-propanoate. A round-bottomed flask filled with 1-(3-methyl-[1,2,4]-triazolo-[4,3-b]pyridazin-6-yl)-hydrazine (190 mg, 1.066 mmol) and glacial acetic acid (1.90 ml) was added methyl 4-acetyl-5-oxohexanoate (0.22 mg, 1.17 mmol). The reaction mixture was stirred at room temperature for 15 min, and then the acetic acid was removed under vacuo. An aqueous solution of 5 % sodium bicarbonate solution was added and the mixture was extracted with ethyl acetate (3x). The combined organic layers were dried over magnesium sulfate, filtered and concentrated in vacuo. The residue obtained was purified by flash chromatography eluting with a gradient of ethyl acetate / methanol $100:00 \rightarrow 90:10$ as eluent. The combination of the appropriate fractions yielded 243mg (66%) of the title compound as a light yellow solid. ${}^{1}H$ **NMR** (400MHz, DMSO-d₆): δ 8.37 (d, J =10 Hz; 1H), 7.85 (d, J = 10 Hz; 1H), 3.6 (s, 3H), 2.73-2.66 (br m, 5H), 2.6 (s, 3H), 2.55-2.48 (br m, 2H), 2.24 (s, 3H); LC-MS (ESI+) m/z: $[M+H]^+$ Calcd. for $C_{15}H_{19}N_6O_2$ 315.35 found 315.40, t_R = 0.98 min.



3-(3,5-Dimethyl-1-(3-methyl-[1,2,4]-triazolo-[4,3-b]-pyridazin-6-yl)-1H-pyrazol-4-yl)propanoic acid. To a solution of methyl 3-(3,5dimethyl-1-(3-methyl-[1,2,4]triazolo[4,3*b*]pyridazin-6-yl)-1*H*-pyrazol-4-yl)propanoate (175 mg, 0.557 mmol) in THF (1.75 ml) and water (0.7 ml) was added LiOH.H₂O (70.16 mg, 1.671 mmol). The reaction mixture was stirred at room temperature for 20 h. A 3N HCl solution (8 ml) was added and the mixture was extracted with a mixture of 20 % isopropanol in dichloromethane (3x). The combined organic layers were dried over magnesium sulfate, filtered and concentrated in vacuo. The white solid obtained (140 mg, 84%) was used without further purification. ¹H NMR (400MHz, DMSO d_6): δ 12.19 (s, 1H), 8.36 (d, J = 10 Hz; 1H), 7.85 (d, J = 10 Hz; 1H), 2.76-2.66 (br m, 5H), 2.6 (s, 3H), 2.55-2.48 (br m, 2H), 2.24 (s, 3H); **LC-MS** (ESI+) m/z: $[M+H]^+$ Calcd. for $C_{14}H_{17}N_6O_2$ 301.32 found 301.40, $t_R = 0.84$ min.



1-(4-Benzylpiperidin-1-yl)-3-(3,5-dimethyl-1-(3-methyl-[1,2,4]-triazolo-[4,3-b]-pyridazin-6yl)-1H-pyrazol-4-yl)-propan-1-one. A roundbottomed flask filled with a solution of 3-(3,5-Dimethyl-1-(3-methyl-[1,2,4]-triazolo-[4,3-*b*]pyridazin-6-yl)-1H-pyrazol-4-yl)-propanoic acid (50 mg, 0.17 mmol), in dimethylformamide (3 ml), were added 4-benzylpiperidine (0.06 ml, 0.33 mmol), HATU (68.52 mg, 0.18 mmol)) and triethyl amine (0.04 ml, 0.306 mmol). The resulting reaction mixture was stirred at room temperature until complete consumption of the starting material (17h) was observed. It was then diluted with water (15 ml) and extracted with ethyl acetate (3x10mL). The combined organic layers were dried over MgSO₄, filtered and concentrated in vacuo. The product was purified by flash chromatography eluting with a linear gradient of ethyl acetate / methanol $100:00 \rightarrow 50:50$ as eluent. The combination of the appropriate fractions yielded 55 mg (70%) of the title product: ¹H NMR (400MHz, DMSO-d_6): δ 8.4 (d, J = 10Hz, 1H), 7.88 (d, J = 10Hz, 1H), 7.33-7.09 (br m, 3H), 7.10-6.99 (br m, 2H), 4.39 (br d, 1H), 3.76 (br d, 1H), 2.86 (t, 1H),

2.71-2.61 (br m, 6H), 2.59 (s, 3H), 2.47-2.14 (br m, 4H), 2.24 (s, 3H), 1.77-1.63 (br m, 1H), 1.59-1.43 (br m, 2H), 0.99-0.73 (br m, 2H); **LC-MS** (ESI+) m/z: $[M+H]^+$ Calcd. for C₂₆H₃₂N₇O 458.58 found 458.61, t_R = 1.16 min.

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Figure 1. C25-140 binds TRAF6, inhibits TRAF6-Ubc13 interaction and TRAF6 activity. *A*, Chemical structure of C25-140 is indicated. *B*, Ectopically expressed HA-TRAF6 RZ₁₂ in HEK-293T cells co-immunoprecipitates less endogenous Ubc13 after C25-140 treatment. Ubc13 levels in the IP fractions were densitometrically quantified in relation to precipitated TRAF6. Error bars indicate mean +/- S.D.; n = 4 biological replicates were quantified; unpaired t-test (two-tailed); ****P<0.0001; D = DMSO. *C*, Heteronuclear correlation NMR experiments using 15N-labeled wild-type TRAF6 RZ₁ either alone (blue) or in the presence of C25-140 (red; molar ratio TRAF6:C25-140 = 1:5). Several NMR signals experience intensity reduction or broadening beyond detection (arrows) indicating binding in so called "intermediate exchange regime". This suggests that protein-compound complex lifetime is in millisecond time scales and confirms direct binding of the compound to TRAF6. *D*, C25-140 dose-dependently counteracts *in vitro* ubiquitination by untagged Wild-type TRAF6 RZ₁ together with the E2 complex Ubc13/Uev1a.



Figure 2. Effect of C25-140 on various E3 ligases/E2 enzyme reactions. *A*, cIAP1 generating K63-ubiquitin chains was also inhibited by C25-140. *B*, MDM2 and TRIM63 generating K48-ubiquitin chains were not impaired by C25-140. *C*, RNF4, a SUMO targeted ubiquitin ligase, was also not affected by C25-140. *D*, Activity of the HECT E3 ligases ITCH and E6AP was not inhibited by C25-140. The substrates and the compound concentrations are indicated; assays were purchased from Boston Biochem.



Figure 3. Effect of C25-140 on E1 and E2 enzymes. Various E1/E2 ubiquitination reactions including UbcH13/Uev1a were carried out in the absence or presence of C25-140 (30 μ M). None of the reactions is affected by the compound. C = control without ATP; D = DMSO; 140 = compound C25-140



Figure 4. C25-140 effect on proinflammatory signaling and T-cell activation. *A*, Endogenous TRAF6 auto-ubiquitination in MEF cells upon IL-1β stimulation is reduced after C25-140 treatment. *B*, C25-140 impairs IL-1β-induced IκBα phosphorylation. pIκBα levels were densitometrically quantified in relation to β-Actin. Error bars indicate mean +/- S.D.; n = 3 biological replicates were quantified; unpaired t-test (two-tailed); ****P<0.0001. *C*, Target gene (ICAM-1 and A20) expression is diminished after IL-1β stimulation and C25-140 treatment; error bars indicate mean +/- S.D.; n ≥ 3 biological replicates; unpaired t-test (two-tailed). *P<0.05, **P<0.01, ****P<0.0001. *D*, Endogenous TRAF6 auto-ubiquitination after PMA/Ionomycin (P/I) stimulation is reduced after C25-140 treatment. *E*, C25-140 decreases IκBα phosphorylation after P/I stimulation in Jurkat T-cells. pIκBα levels were densitometrically quantified in relation to β-Actin. Error bars indicate mean +/- S.D.; n = 3 biological replicates were quantified; unpaired t-test (two-tailed); *P<0.05, **P<0.01. *F*, Upon P/I stimulation, IL-2 and TNFα cytokine secretion is attenuated after C25-140 treatment. Error bars indicate mean +/- S.D.; n = 3 biological replicates; unpaired t-test (two-tailed); *P<0.05, **P<0.01. *F*, Upon P/I stimulation, IL-2 and TNFα cytokine secretion is attenuated after C25-140 treatment. Error bars indicate mean +/- S.D.; n = 3 biological replicates; unpaired t-test (two-tailed); *P<0.05, **P<0.01; D = DMSO.



Figure 5. C25-140 impedes immune receptor signaling in primary human PBMC and primary murine T-cells. *A-C*, PBMCs from healthy individuals were isolated and treated with C25-140 prior to stimulation with *A*, IL-1 β , *B*, LPS and *C*, CD3/CD28. Secretion of various NF- κ B driven cytokines was measured by ELISA and showed reduction of distinct cytokine secretion by C25-140 in all donors. *D*, Primary mouse CD4+ T-cells were isolated and treated with C25-140 and subsequently stimulated with CD3/CD28. IL-2 mRNA expression and protein secretion were measured using qRT-PCR and ELISA, respectively. C25-140 dose-dependently reduced IL-2 expression; error bars indicate mean +/- S.D.; n = 4 biological replicates; signals were normalized to DMSO; unpaired t-test (two-tailed); **P<0.01, ***P<0.001, ****P<0.0001; D = DMSO.



Figure 6. Topical application of the inhibitor C25-140 reduces *in vivo* outcome of imiquimodinduced psoriasis. *A*, Preclinical mouse model for imiquimod (IMQ)-induced psoriasis. Mice (n = 8 per group) were topically treated with IMQ to the back and ear once a day. C25-140 was topically applied twice daily. Mice were scored throughout the study and samples were collected at day 6 for cytokine measurement. (B-E) Mice were scored for *B*, cumulative score, *C*, thickness score, *D*, scaling and *E*, erythema. All symptoms were improved by C25-140 treatment; error bars indicate mean +/- S.D.; 2-way ANOVA test (Sidak's multiple comparison test). *F*, IL-17 cytokine levels were evaluated by ELISA and show a significant decrease after treatment; Unpaired t-test (two-tailed); error bars indicate mean +/- S.D.; *P<0.05, **P<0.01, ***P<0.001, ***P<0.0001.



Figure 7. C25-140 ameliorates symptoms of rheumatoid arthritis (RA) in a preclinical mouse model. *A*, Study design of a collagen-induced arthritis (CIA) preclinical mouse model for RA (n = 10 per group). RA was induced by collagen application at day 0 and a booster injection at day 21. At day 28, C25-140 was intraperitoneally applied twice daily for 14 days. Mice were scored throughout the study and limbs were collected at day 14 of treatment for histopathology. *B*, The average arthritic index was determined for each group to analyze efficacy of treatment. C25-140 and the control prednisolone significantly reduced the arthritic index; error bars indicate mean +/- S.E.M.; 2-way ANOVA test (Sidak's multiple comparison test). In histopathology analyses, mice were scored for *C*, H&E staining of sections for histopathology (ankle and hindpaw). C25-140 at 14 mg/kg ameliorates disease outcome; more histology images in Figure S7. *D*, Summed histology score, *E*, inflammation, pannus, and cartilage damage. All quantified parameters demonstrate a dose-dependent improvement of RA symptoms after C25-140 treatment; error bars indicate mean +/- S.E.M.; One-way ANOVA test (Sidak's multiple comparison test); more quantification of histology data are in Figure S8; ns = not significant; ***P<0.001, ****P<0.0001; BID = twice daily.

Targeting TRAF6 E3 ligase activity with a small molecule inhibitor combats autoimmunity

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