Production of biologically active IL-36 family cytokines through insertion of N-terminal caspase cleavage motifs

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Recent evidence has strongly implicated IL-36 cytokines as key initiators of inflammation in the skin barrier. IL-36 cytokines belong to the extended IL-1 family and, similar to most members of this family, are expressed as inactive precursors that require proteolytic processing for activation. Because the proteases responsible for activation of members of the IL-36 subfamily have not been reported, we have developed a method for the production of biologically active IL-36 through introduction of a caspase cleavage motif, DEVD, within the N-termini of these cytokines. Here, we show that DEVD-modified IL-36α, IL-36β and IL-36γ cytokines were highly soluble and were readily processed and activated by caspase-3. Caspase-3-processed IL-36 family cytokines exhibited robust biological activity on a range of responsive cell types, including primary keratinocytes. We also generated specific polyclonal antibodies against all three IL-36 family members through immunization with purified recombinant IL-36 cytokines. The modified forms of IL-36 described herein will be useful for production of large quantities of biologically active IL-36 for structure and function studies on these important proinflammatory cytokines.

IL-1 family cytokines, which include the recently described IL-36α, IL-36β and IL-36γ proteins, play major roles as initiators of inflammation and are frequently among the first cytokines produced in response to infection or injury [1–4]. IL-1 family cytokines can initiate complex cascades of additional cytokine production from diverse cell types [5–10]. IL-36α, IL-36β and IL-36γ are encoded by distinct genes and accumulating evidence suggests that these cytokines play a key role in skin inflammation, particularly in psoriasis [11–20]. Members of the IL-1 family are typically among the most upstream cytokines to be released upon infection or cell damage and most likely represent the key damage-associated molecular patterns (DAMPs) as a consequence of their release during necrosis [4,21].

IL-36α, IL-36β and IL-36γ are all generated as leaderless cytokines that lack biological activity and are not secreted via the canonical endoplasmic reticulum (ER)–golgi secretion route [22]. Thus, proteolytic processing of IL-36 cytokines is required to unleash their proinflammatory activity, similar to other members of the IL-1 family, such as IL-1β and IL-18 [23,24]. Sims and colleagues have shown that removal of a small number of residues from the N-termini of IL-36α, IL-36β and IL-36γ increases their biological activity by greater than 10 000-fold [22]. However, the proteases

Abbreviations
Ac-DEVD-AFC, Ac-Asp-Glu-Val-Asp-7-amino-4-trifluoromethylcoumarin; DEVD, Asp-Glu-Val-Asp; DMEM, dulbecco’s modified eagle’s medium; ER, endoplasmic reticulum; GST, glutathione S-transferase; IL-1AcP, IL-1 receptor accessory protein; IL-36RA, IL-36 receptor antagonist; IL-36R, IL-36 receptor; IPTG, isopropyl β-D-1-thiogalactopyranoside; NFDM, non-fat dry milk; Ni-NTA, nickel-nitrilotriacetic acid; PARP, poly (ADP-ribose) polymerase; PMA, phorbol 12-myristate 13-acetate; poly(I:C), polyinosinic:polycytidylic acid; PRB, protease reaction buffer; RFU, relative fluorescence units; SEAP, secreted embryonic alkaline phosphatase; SUMO, small ubiquitin-like modifier; TEV, tobacco etch virus; Ulp1, ubiquitin-like-specific protease 1.
responsible for activation of IL-36 cytokines are currently unknown.

To facilitate investigations into the biological activity of the IL-36 cytokine subfamily, a simple and efficient approach for the production of biologically active IL-36 cytokines is desirable. Here, we describe a novel strategy for the production of large amounts of soluble active recombinant IL-36 cytokines through insertion of a caspase-3 cleavage motif (DEVD) within the N-terminals of these cytokines, followed by processing by purified recombinant caspase-3. This approach enabled us to process and activate human IL-36 cytokines with exquisite specificity, leading to a dramatic increase in biological activity. Here, we show that caspase-3-processed recombinant IL-36 is highly active on a range of cell types, including primary keratinocytes, and elicits robust production of multiple cytokines and chemokines from target cells. We also established a convenient reporter cell line HeLaIL-36R overexpressing the IL-36 receptor, as well as an NFκB-SEAP reporter plasmid, which will be useful as a screening system to identify molecules involved in IL-36 activation and downstream signalling. Using purified forms of IL-36, we also generated polyclonal antibodies against all three IL-36 cytokines that displayed specificity for individual IL-36 proteins.

Results

Generation of DEVD-modified IL-36 cytokines

Previous studies have shown that the full-length forms of IL-36α, IL-36β and IL-36γ exhibit little or no bioactivity [22], similar to other members of the extended IL-1 family, but become active upon removal of small stretches of N-terminal amino acids. For example, truncation of IL-36α after Leu5 has been shown to enhance its activity 10 000-fold, similarly, truncation of IL-36β after Gln4 and IL-36γ after Gln17 also unleashed the activity of the latter cytokines [22]. However, the proteases that naturally process at these sites to activate the latter cytokines have not been reported to date.

To facilitate activation of recombinantly expressed IL-36 cytokines for further study of their biological activity, we generated insertions within the N-terminal region of all three IL-36 coding sequences to insert the known artificial processing sites [22]. This approach permits the generation of highly precise protein truncations to eliminate purification tags, autoinhibitory domains, or other unwanted protein domains prior to use in biological assays.

The DEVD motifs were inserted at regions within IL-36 cytokines previously identified by Sims and colleagues to result in the liberation of robust biological activity upon proteolysis at these sites [22]. We then transformed BL-21 (RIL) strain E. coli with pET45b-based IL-36 expression plasmids, followed by induction of protein expression for 3 h at 37 °C. As shown in Fig. 2, all three IL-36 cytokines were strongly expressed under these conditions and were readily purified as soluble proteins from the supernatant fraction of sonicated bacterial cell lysates. Proteins were washed and eluted from nickel-NTA affinity matrix and yields were estimated at ~10 μg·mL⁻¹ of bacterial culture.

Processing of DEVD-modified IL-36 by purified caspase-3

To facilitate processing of DEVD-modified IL-36 cytokines, we expressed and purified recombinant
caspase-3 as previously reported [25]. Purified caspase-3 was highly active as demonstrated by its ability to cleave the synthetic caspase substrate DEVD-AFC (Fig. 3A). We next incubated DEVD-modified IL-36α, β and γ with recombinant caspase-3 over a range of concentrations from 1 μM to 200 nM and, as shown in Fig. 3B, C, this resulted in cleavage of the DEVD-modified form of IL-36β (DEVDIL-36β) to a slightly smaller product consistent with removal of the N-terminal residues as desired. Similar caspase-3 cleavage reactions were also conducted with the DEVD-modified forms of IL-36α and IL-36γ, with identical results (data not shown).

DEVD-modified IL-36 cytokines are biologically active

To ask whether caspase-3-cleaved DEVD-modified IL-36 cytokines were biologically active, we titrated full-length versus caspase-3-cleaved forms of DEVDIL-36α, β and γ onto HeLa cells that were stably transfected with the human IL-36 receptor. As shown in Fig. 4A, whereas all three full-length DEVDIL-36 cytokines were completely inactive in their unprocessed forms, processing of DEVDIL-36 cytokines with caspase-3 resulted in highly robust biological activity as demonstrated by the production of very high levels of IL-6, IL-8 and CXCL1 by HeLaIL-36R cells. Importantly, caspase-3 alone had no effect on cytokine production by HeLa cells (Fig. 4B), excluding the unlikely possibility that addition of the latter protease to the IL-36 preparations could be the source of increased cytokine activity. We also examined some of the intracellular signal transduction events seen in response to active IL-36. As shown in Fig. 4C, treatment of HeLaIL-36R cells with biologically active IL-36 resulted in the rapid phosphorylation and proteolytic processing of the NFκB inhibitor IκB, in tandem with phosphorylation of the p65 subunit of NFκB, as well as phosphorylation of p38MAPK. Collectively, the above data clearly demonstrate that removal of the N-terminal region of IL-36 cytokines through insertion of a tetrapeptide caspase cleavage motif permitted the simultaneous removal of the poly-histidine purification tag as well as
the N-terminal region responsible for autoinhibition of IL-36 family cytokines, leading to the generation of highly active IL-36.

**Generation of a reporter cell line for the assessment of IL-36 activity**

Because IL-36 promotes NFκB activation (Fig. 4C), we also generated a convenient reporter cell line, based upon the NFκB-dependent expression of secreted alkaline phosphatase (SEAP), to assess IL-36 activity. As shown in Fig. 5A, HeLa-IL-36R cells stably transfected with the NFκB-SEAP reporter exhibited robust SEAP activity upon treatment with caspase-3-cleaved DEVD-IL-36. As expected, HeLa-IL-36R-SEAP cells also secreted multiple cytokines in response to IL-36 treatment (Fig. 5B). This reporter line will be useful for future studies aimed at identifying upstream components of IL-36 signal transduction events and also for screening of inhibitors of IL-36 signalling.

**Transformed and primary keratinocytes are responsive to IL-36**

Because the preceding tests of the biological activity of IL-36 cytokines were conducted on cells overexpressing the IL-36 receptor, we also asked whether cells that naturally express the IL-36R were IL-36 responsive. As noted in the Introduction, IL-36 is strongly linked with psoriasis which is thought to result from excessive inflammation in the epidermal layers of skin, driven in part through the release of inflammatory mediators from keratinocytes. Thus, we asked whether transformed HaCaT keratinocytes, as well as primary human keratinocytes were responsive to caspase-3-cleaved DEVD-modified IL-36 cytokines. As shown in Fig. 6, both cell types were highly responsive to caspase-3-cleaved, but not full-length, DEVD-IL-36, once again confirming that the former is highly active and can be readily produced in large amounts for the exploration of IL-36 function in cellular and molecular studies. Furthermore, these data also show that insertion of a simple tetrapeptide caspase cleavage motif is a useful strategy for the facile removal of purification tags or protein domains that interfere with protein solubility or activity.

**Generation of polyclonal antibodies against individual IL-36 proteins**

Finally, we also used recombinant IL-36 proteins to immunize rabbits with individual IL-36 proteins (i.e. α, β or γ) to generate polyclonal antibodies capable of recognizing the latter cytokines (Fig. 7A). As Fig. 7B illustrates, this approach yielded antibodies that were highly specific for IL-36β and IL-36γ, with little cross-reactivity against each other, as well as antibodies that were relatively specific for IL-36α. Moreover, the IL-36γ polyclonal antibody was capable of detecting endogenous levels of IL-36γ from primary human keratinocytes (Fig. 7C). These antibodies will be useful for investigations exploring the factors that influence IL-36 expression in healthy versus inflamed tissues.

**Discussion**

Here, we have shown that the insertion of a caspase-3 cleavage motif (DEVD) into IL-36 cytokines enabled us to produce highly active and soluble forms of these cytokines upon processing with caspase-3. These data also confirmed that proteolytic processing of IL-36α, β and γ dramatically increase the biological activity of IL-36.
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these cytokines. We also established a HeLa IL-36R bioassay to screen for proteases that naturally activate IL-36 cytokines.

IL-36 cytokines require N-terminal processing for activation, however the proteases responsible for this activation are unknown. Thus, the insertion of DEVD motifs allowed us to potently activate IL-36 cytokines using caspase-3 to generate truncations at sites previously identified to liberate biological activity within these cytokines [22]. Caspases have exquisite specificity for cleavage after aspartic acid residues and so can selectively and efficiently process the DEVD tetrapeptide which is not commonly found in many proteins, especially within surface exposed loop regions prone to proteolysis. The DEVD peptide was originally identified from the caspase cleavage site within PARP, a DNA repair enzyme, targeted by caspase-3 during apoptosis [27]. Derivatives of DEVD are frequently used to assess caspase-3 enzymatic activity by fluorimetric methods and also as inhibitors of caspase-3 activity. Here, we have demonstrated a novel use of the DEVD motif as a fusion tag to generate proteins that are susceptible to proteolysis by caspase-3. Similar strategies have been previously used by other groups to purify biologically active forms of IL-36RA and IL-38 [22,28]. IL-36RA requires removal of the N-terminal Met1 residue to act as an antagonist of IL-36 signalling but exhibits little activity when this residue is present [22]. Towne et al. used a factor Xa protease removable N-terminal GST tag to generate truncated IL-36RA. However, one potential drawback of the latter approach is that GST, being a relatively large fusion partner (26 kDa), may introduce solubility and folding problems in proteins tagged using this strategy. Furthermore, removal of the GST tag by factor Xa has been reported to be relatively ineffective and nonspecific, reducing yields of the desired cleavage product [29,30].

Small ubiquitin-related modifier (SUMO) fusion tags have also been used to purify N-terminally truncated...
IL-36 ligands [22]. SUMO is a relatively small protein tag (~11 kDa) often used as a solubility enhancer for protein purification by promoting proper folding and stability. After purification, the SUMO moiety can be removed by the highly specific SUMO protease UlpI. However, unlike the GST tag, SUMO requires an additional affinity tag for purification of the fusion protein, usually a polyhistidine tag [30]. Hu et al. recently described the purification of IL-36 as a TEV fusion protein. TEV protease, from the tobacco etch virus, is more stringent than factor Xa and other commonly used fusion tag proteases like thrombin and enterokinase. Wild-type TEV protease can be deactivated by autolysis and expression of the protease is hindered by solubility issues, however, mutated forms of the protease have been developed for increased stabilization and expression yields [31]. Inclusion of the small caspase-3-specific peptide tag DEVD acts as a simple and reliable alternative for accurate removal of purification tags and inhibitory domains within proteins with minimal disruption to the fusion partner.

We also established a convenient reporter cell line HeLaIL-36R, which overexpresses the NFκB-responsive SEAP reporter plasmid. This bioassay permits the rapid screening of compounds that naturally antagonize IL-36 cytokine activity. Activation of NFκB signalling by IL-36 induces the release of secreted alkaline phosphatase into culture medium that is highly stable and can be readily detected by colorimetric assay. This allows the rapid and continuous moni-

Fig. 6. Truncated IL-36 proteins exhibit biological activity on primary and transformed keratinocytes. (A) DEVDIL-36b (250 nM) was incubated with caspase-3 (400 nM) for 2 h at 37 °C. HaCaT cells (5 x 10⁴ cells per well) were stimulated with DEVDIL-36b, as indicated. After 18 h, cytokine concentrations in the culture supernatants were determined by ELISA. (B) Primary human keratinocytes (4 x 10⁴ cells per well) were stimulated with DEVDIL-36b, as indicated. After 18 h, cytokine concentrations in the supernatants were determined by ELISA. Error bars represent the mean ± SEM of duplicate determinations from a representative experiment.
Monitoring of NF-κB activation without the need to disrupt or lyse cell populations.

It is interesting that removal of very small numbers of residues from the N-termini of IL-36 family cytokines have such dramatic effects on the biological activities of these cytokines. For example, the IL-36 N-termini may partly occlude the receptor binding domain through steric clashes, removal of which permits a more stable interaction with the IL-36R complex. Alternatively, proteolysis of IL-36 cytokines may induce a conformational change in these proteins that increases their affinity for the IL-36 receptor, thereby increasing biological potency. Interestingly, Hazuda and colleagues have reported that IL-1α and IL-1β cytokines undergo profound conformational changes upon removal of their N-termini and, as a consequence, the mature regions of these molecules switch from a proteinase K-sensitive to a proteinase K-insensitive state [32]. This change is most likely reflected in an altered conformation that increases affinity for the IL-1 receptor [32]. The further resolution of IL-36 ligand structures, both in their full-length, as well as their processed forms by naturally occurring proteases, will help to elucidate how exactly these molecules interact with the IL-36R/IL-1AcP receptor complex.

In conclusion, here we have described a novel method to produce highly soluble and active recombinant IL-36 cytokines through insertion of a small caspase cleavage motif within the N-terminal region of these cytokines, followed by processing by caspase-3. IL-36 cytokines have been reported to have an important role in the pathogenesis of psoriasis, a common skin inflammatory condition. We have shown both transformed and primary keratinocytes, which naturally express IL-36R, were highly responsive to active caspase-3-cleaved DEVD-modified IL-36. This method facilitates production of active IL-36α, IL-36β and IL-36γ that can be used to further investigate the role of the IL-36 subfamily in inflammation.

**Experimental procedures**

**Materials**

Anti-IL-36β antibody was purchased from R&D (Minneapolis, MN, USA) systems. Antibodies specific to p-p65, p-p38 and p38 were obtained from Cell Signalling Technology (Danvers, MA, USA). Anti-p65 and anti-IκBα were from Santa Cruz Biotechnology (Dallas, TX, USA). Anti-p-p38 and p38 were obtained from Cell Signalling Technology (Danvers, MA, USA). Anti-IL-36 antibodies described in (B). Recombinant IL-36 (500 pg) serves as a positive control for each immunoblot.

**Cell culture**

The HeLaIL-36R cell line was generated by transfection with pCXN2-IL-1Rrp2 (IL-36R) plasmid followed by selection using G418 antibiotic (Sigma). IL-36R overexpressing clones were expanded from single cells using limiting dilution cloning, followed by expansion of individual clones. The HeLaIL-36R-SEAP cell line was generated by transfection of stable HeLaIL-36R overexpressing cells with pNifty2-SEAP plasmid (InvivoGen) followed by selection using Zeocin antibiotic. Clones were expanded from single cells using limiting dilution cloning. Clones were selected by demonstration of acquired optimal responsiveness to active forms of IL-36 via SEAP production and ELISA. HaCaT cells were cultured in Dulbecco’s modified eagle’s medium (DMEM) (Gibco,
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Waltham, MA, USA) supplemented with FCS (10%). Primary neonatal foreskin-derived keratinocytes were purchased from Cell Systems (Troisdorf, Germany) and cultured in serum-free Dermalife K media (Cell Systems). All cells were cultured at 37 °C in a humidified atmosphere with 5% CO₂.

Expression and purification of recombinant DEVDIL-36 and caspase-3

Modified forms of IL-36 containing a caspase-3-processing motif (DEVD) N-terminal to known processing sites [22] were generated by cloning the DEVD-modified cytokine coding sequence in frame with the poly-histidine tag sequence in the bacterial expression vector pET45b. Proteins were expressed by addition of 600 µM IPTG to exponentially growing cultures of BL21-CodonPlus(DE3)-RIL bacteria followed by incubation for 3 h at 37 °C. Bacteria were lysed by sonication and poly-histidine tagged proteins were captured using nickel-NTA agarose (Amintra, Expedeon Ltd, Cambridgeshire, UK), followed by elution into PBS, pH 7.2, in the presence of 100 mM imidazole. Recombinant poly-histidine-tagged caspase-3 was expressed and purified as described previously [25]. Bacterial cell lysates and purified proteins were visualized on 12% SDS/PAGE electrophoresis gel stained with Coomassie Brilliant Blue.

Caspase-3 activity assay

Protease cleavage reactions (40 µL final volume) were carried out in protease reaction buffer (50 mM HEPES, pH 7.4, 75 mM NaCl, 0.1% CHAPS, 2 mM DTT) containing 50 µM Ac-DEVD-AFC. Samples were measured by using an automated fluorimeter (Spectrafluor Plus; TECAN, Reading, UK) at wavelengths of 430 nm (excitation) and 535 nm (emission) for 60 min.

Caspase-3 cleavage assays

Reactions were carried out in protease reaction buffer (50 mM HEPES, pH 7.4, 75 mM NaCl, 0.1% CHAPS, 2 mM DTT) for 2 h at 37 °C. Full-length or processed DEVDIL-36 cytokines were then used to treat cells (5 x 10⁴ cells per well) for 18–24 h, followed by assessment of cytokine production by ELISA.

Measurement of cytokines and chemokines by ELISA

Cells were plated at 5 x 10⁴ cells per well and treated with various forms of IL-36 as indicated in the appropriate figure legends. Cytokines and chemokines were measured from cell culture supernatants using specific ELISA kits obtained from R&D Systems (human IL-6, IL-8, CXCL1, CCL20). All cytokine assays were carried out using duplicate samples from each culture.

Measurement of NFκB activation by SEAP assay

HelaIL36R-SEAP cells were plated at 5 x 10⁴ cells/well in a tissue culture dish in RPMI supplemented with 5% heat-inactivated FCS. Cells were treated with a titration of full-length or caspase-3-activated DEVDIL-36β. Cell culture supernatants were harvested and secreted alkaline phosphatase in the medium was detected using Quantiblue reagent (InvivoGen). Ten microlitres of cell culture supernatant was incubated with 190 µL prewarmed Quantiblu reagent and incubated for 15 min at 37 °C. To assess SEAP activity, absorbance readings were taken at 620 nm with a microplate reader (Infinite F50; TECAN) after 15 min to 6 h incubation periods.

Western immunoblotting

Protein samples were prepared using SDS/PAGE loading buffer (2% SDS, 50 mM Tris–HCl, pH 6.8, 10% glycerol, 2.5% β-mercaptoethanol), boiled for 7 min and electrophoresed on 12% SDS/PAGE gels. Proteins were then transferred onto nitrocellulose membrane at 40 mA overnight. Membranes were blocked for 1 h (5% NFDM, 0.05% sodium azide in Tris-buffered saline, Tween-20, TBST). The indicated proteins were probed using specific antibodies, typically diluted 1:1000. Membranes were washed 3 times in TBST and then incubated with the relevant HRP-conjugated secondary antibody diluted 1:2000. Membranes were again washed and proteins were visualized with SuperSignal West Pico (Thermo Scientific, Waltham, MA, USA) and exposure to autoradiography films.

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Author contributions

D.M.C. and C.M.H. performed experiments, analysed data, generated the figure panels and wrote the figure legends. G.P.S., P.B.D. and E.B. purified recombinant proteins and validated some of the cell line data. S.J.M conceived the study, designed and anal-
ysised experiments, supervised the study and wrote the manuscript with contributions from D.M.C. and C.M.H.

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