Cold-induced urticarial autoinflammatory syndrome related to factor XII activation

Jörg Scheffel1,2,6, Niklas A. Mahnke1,2,6, Zonne L.M. Hofman3, Steven de Maat3, Jim Wu1, Hanna Bonnekoh1,2, Reuben J. Pengelly4, Sarah Ennis4, John W. Holloway4, Marieluise Kirchner5, Philipp Mertins5, Martin K. Church1, Marcus Maurer1,2*, Coen Maas3 & Karoline Krause1,2

Hereditary autoinflammatory diseases are caused by gene mutations of the innate immune pathway, e.g. nucleotide receptor protein 3 (NLRP3). Here, we report a four-generation family with cold-induced urticarial rash, arthralgia, chills, headache and malaise associated with an autosomal-dominant inheritance. Genetic studies identify a substitution mutation in gene F12 (T859A, resulting in p.W268R) which encodes coagulation factor XII (FXII). Functional analysis reveals enhanced autocatalytic cleavage of the mutated protein and spontaneous FXII activation in patient plasma and in supernatant of transfected HEK293 cells expressing recombinant W268R-mutated proteins. Furthermore, we observe reduced plasma prekallikrein, cleaved high molecular weight kininogen and elevated plasma bradykinin. Neutrophils are identified as a local source of FXII. Interleukin-1β (IL-1β) is upregulated in lesional skin and mononuclear donor cells exposed to recombinant mutant proteins. Treatment with icatibant (bradykinin-B2-antagonist) or anakinra (interleukin-1-antagonist) reduces disease activity in patients. In conclusion, our findings provide a link between contact system activation and cytokine-mediated inflammation.
Systemic autoinflammatory diseases (SAIDs) are characterized by self-directed inflammation that is mediated via activation of innate immune pathways. The majority of SAIDs are rare and disabling disorders with significant morbidity and quality of life impairment. Within the last 15 years, the molecular basis of several hereditary SAIDs was uncovered by using novel high-throughput sequencing techniques. Many of the affected genes were shown to involve molecules associated with inflammasome signaling. Inflammasomes are intracellular multiprotein complexes, which assemble upon recognition of diverse pathogen- and danger-associated molecular patterns (PAMPs/DAMPs) leading to activation of caspase-1 to convert pro-interleukin-1β (pro-IL-1β) and pro-IL-18 into their active forms. Consequently, cytokine-targeted therapies, in particular directed against IL-1β and its receptor, largely improved the treatment of SAID patients.

Cryopyrin-associated periodic syndrome (CAPS) is one of the best-studied hereditary SAIDs. Its hallmark feature is recurrent urticarial rash combined with systemic symptoms, such as fever episodes, arthropathy, myalgia, headaches, and malaise. Cold temperatures represent a major trigger of the disease. CAPS is caused by a heterozygous mutation in the gene encoding for nucleotide binding like receptor protein 3 (NLRP3 or cryopyrin)1. The NLRP3 mutation in CAPS results in an overactivated inflammasome followed by IL-1β-mediated inflammation. A diagnosis of CAPS is often delayed, and only eventually made by excluding other immune-mediated disorders, such as infections, immunodeficiencies, autoimmune, and neoplastic diseases. Previously, up to 40% of cases with typical CAPS phenotype were reported to lack disease-causing NLRP3 mutations, suggesting the existence of other, as of yet unknown genetic variants2. The increased use of sequencing techniques identified a number of somatic CAPS mutations in patients tested negative by conventional Sanger sequencing (depending on the study and phenotype in 12–69% of individuals)3,4. Besides CAPS, few other genetic defects were found, including mutations in NLRP12, NLRC4, and PLCG2. All of them present with early onset urticarial rash triggered by cold and an autoinflammatory phenotype5–9.

Here, we describe an autosomal-dominant disease characterized by cold-induced urticarial rash and systemic inflammation. The disease is associated with a mutation in the human coagulation Factor 12 gene (F12), and we propose the term FXII-associated cold autoinflammatory syndrome (FACAS) to denote this disease. Functional studies show autocatalytic cleavage of FXII and profound activation of the contact system linked to IL-1β-mediated inflammation.

Results
Clinical phenotype and disease manifestation. Four family members of a four-generation family of European ancestry presented with cold-induced urticarial rash that started within the first weeks after birth (Table 1; Fig. 1a). The non-pruritic burning rash occurred on a daily basis at the extremities, face, and trunk, and was not accompanied by angioedema. Wheal development typically appeared within 10–30 min after whole-body exposure to cold temperatures (<15–20 °C) and stayed for one-to-several hours after rewarming. The urticarial rash was exacerbated by windy and humid weather. Wheals were induced by sweating and consecutive evaporative cooling. Direct localized skin contact with cold fluids or objects, however, did not cause any symptoms. The skin symptoms associated with signs of systemic inflammation including chills, joint pain, headache, fatigue, and subfebrile evening temperatures. All patients had perennial symptoms that deteriorated during the cold season and frequently lead to absence from school and work. Over the course of their disease, patients reported worsening of systemic manifestations, e.g., joint pain, fatigue, and chills, resulting in a more severe phenotype in adult patients. The clinical signs were combined with mild-to-moderate increases in serum levels of the inflammation markers amyloid A (SAA) and S100 A12/S100 A8/9, but not C-reactive protein (CRP). Other routine laboratory markers (e.g., liver and kidney function, blood count, coagulation) were unremarkable. Histopathology of lesional skin following cold exposure revealed moderate dermal edema and perivascular infiltrates composed of macrophages and few neutrophils (Fig. 1b). In contrast to acquired cold urticaria, affected individuals presented with generalized urticarial rash upon cold exposure and tested negative in the ice cube skin provocation test. Cold water bath provocation of the forearm (15 °C for 10 min) did also not cause any wheals or angioedema. In addition, all patients reported complete unresponsiveness to oral antihistamines at standard or higher than standard doses (please see Supplementary Table 1 for a comparison of FACAS with other cold-induced urticarial disorders). A therapeutic trial with IL-1 receptor antagonist anakinra in patients #1 and #2 resulted only in limited (#2) to moderate improvement (#1) of clinical symptoms. However, continuous anti-IL-1 treatment could not be maintained due to severe injection site reactions.

Genetic analysis. Based on the clinical phenotype resembling CAPS, Sanger sequencing of NLRP3 and NLRP12 was initially performed, but failed to identify plausible pathogenic mutations in the index patient (#1). The following whole-exome sequencing prioritized the assessment of genes that were previously associated with a skin phenotype of urticaria or urticaria-like symptoms as reported by the Human Gene Mutation Database (Release 2015.4)10. These were TBX2A2, CEP68, F12, SERPING1, NLRP3, NLRP12, KIT, IL18, HNMT, FCER1α, NLRCA4, ADGRE2, and PLCG2. Sequencing data were generated for three affected family members across three generations (#2, #5, #7, Fig. 1c) and filtered to identify a single candidate variant, a heterozygous substitution mutation in exon 9 of F12 (Fig. 1d). The gene encoding for the human coagulation factor XII (NM_000505.5.c: T859A:p.W268R). The observed variant is not listed in the Genome Aggregation Consortium (gnomAD) database listing ~140,000 individuals11. In silico predictors of variant pathogenicity concordantly inferred deleterious impact (GERP = 5.01; SIFT = 0.0; PolyPhen2 = 1.0). Sanger sequencing of the available family members confirmed consistent segregation of this mutation with the prior annotation of disease status (Fig. 1c, d).

FXII exerts diverse functions in coagulation, fibrinolysis, complement activation, and the kallikrein–kinin system. Mutations in F12 are linked to hereditary angioedema with normal C1 inhibitor (FXII-HAE)12. FXII-HAE is characterized by acute attacks of localized tissue swellings that are mediated by the vasoactive peptide bradykinin, which is produced by the plasma kallikrein–kinin system. Known FXII-HAE mutations are primarily located in the proline-rich region on exon 913,14. In contrast, the W268R variant lies within the triple-looped kringle domain of FXII.

Functional analyses. Markers of coagulation (activated partial thromboplastin time [aPTT], fibrinogen, plasminogen, FXI activity, D-dimers, INR) and complement activation (C3, C4, C1-esterase inhibitor [C1-INH] concentration and function) were normal in all affected family members.

To test, if the FXII-p.W268R substitution in our patients leads to cleavage and/or activation of FXII associated with contact system activation and subsequent generation of bradykinin, we
Table 1 Individual clinical characteristics and manifestations in FACAS patients.

| Characteristic                  | Patient ID #1 | Patient ID #2 | Patient ID #5 | Patient ID #7 |
|--------------------------------|---------------|---------------|---------------|---------------|
| Age at diagnosis               | 35            | 64            | 9             | 27            |
| Sex                            | Female        | Male          | Female        | Female        |
| Age at disease onset           | After birth   | After birth   | After birth   | After birth   |
| Skin symptoms                  | Cold-induced urticarial rash | Cold-induced urticarial rash | Cold-induced urticarial rash | Cold-induced urticarial rash |
| CNS symptoms                   | Headache      | Headache      | Headache      | Headache      |
| Musculo-skeletal symptoms      | Moderate arthralgia | Severe arthralgia | None          | Mild arthralgia |
| Systemic symptoms              | Fatigue, subfebrile temperatures | Fatigue, subfebrile temperatures | Fatigue      | Fatigue      |
| $S_{100}$ A12 in ng/ml (ref. <200) | 523.7         | n.a.          | 1024.3        | n.a.          |
| $S_{100}$ A8/9 in µg/ml (ref. <2.94) | n.a.          | 4.10          | n.a.          | 4.51          |
| Serum amyloid A in mg/l (ref. <6.40) | 16.80         | n.a.          | n.a.          | 8.20          |

**Fig. 1** Phenotype and genotype in patients with FXII-associated cold autoinflammatory syndrome (FACAS).  

**a** Urticarial rash of the face of a 9-year-old girl and the leg of her 35-year-old mother.  
**b** Skin histopathology reveals dermal edema (H.E. yellow arrow) and perivascular infiltrates composed of macrophages (CD163) and few intravascular neutrophils (myeloperoxidase [MPO], yellow arrow). Original magnification ×400 (n = 2 technical replicates).  
**c** Family pedigree over four generations identifying affected (shaded black) and unaffected (unshaded) individuals. Exome sequenced individuals are underlined. Where conducted, the results of Sanger sequencing of p.W286R are shown to top left of each individual whereby “+” denotes heterozygote for variant and wild-type, and “−” denotes healthy reference.  
**d** Representative sequencing chromatogram of the heterozygous genotype in an affected individual (top) and chromatogram of healthy individual from the same region (bottom).
first assessed FXII fragmentation and activity. For its activation, FXII needs to be cleaved into a two-chain molecule composed of a heavy- and a light-chain linked by a disulphide bond. In addition to the unprocessed FXII (78 kDa), molecular analyses revealed an increased presence of the 50 kDa band (reducing) in the FXII immunoblot from plasma of affected family members, which was hardly detectable in healthy control family members and an FXII-HAE control (Fig. 2a). This band resembled the heavy chain of active FXII (FXIIa), suggesting partial fragmentation and activation of FXII in affected patients. Accordingly, recombinant FXII-p.W268R (rFXII W268R) expressed in HEK293 cells also showed the characteristic 50 kDa (III) band in the FXII immunoblot resembling the heavy chain. This band is also present in purified aFXIIa, but not found in unprocessed recombinant FXII (rFXII wt) (Fig. 2b). In addition to the heavy chain, the light chain at 25 kDa (IV) was found in preparations of aFXIIa and rFXII W268R, but not in rFXII wt. Treatment of rFXII W268R with kaolin resulted in increased generation of the light-chain fragment. Besides the heavy and light chains at 50 kDa and 25 kDa, respectively, we found a small, rFXII W268R preparation a third fragment with an apparent molecular weight of 60 kDa (reducing) that was absent in the aFXIIa preparation (Fig. 2a, b). Mass spectrometry analysis of excised gel fragments confirmed the autoproteolytic cleavage of rFXII W268R at residues AA353/354, which yields in FXIIa. In addition, analysis of the 60-kDa fragment revealed cleavage of rFXII W268R after AA447, which is part of the protease domain and facilitated by kaolin treatment (Fig. 2c). In line with these observations which suggest autoactivation of FXII W268R, we found high spontaneous activity of FXIIa in the plasma of affected family members (Fig. 2d) as well as in the supernatant of FXII W268R-expressing HEK293 cells, but not in the plasma of healthy family members or wt-FXII expressing HEK293 cells (Fig. 2d, e). Interestingly, overall plasma levels of FXII were comparable with healthy controls (Supplementary Fig. 1), indicating that the p.W268R mutation only affects the probability of spontaneous FXII activation, but not its expression.

We next assessed mediators of the kallikrein–kinin pathway. High FXIIa activity was associated with pronounced plasma prekallikrein (PK) cleavage and kallikrein formation (Fig. 3a) similar to what is found in activated recombinant PK preparations (Supplementary Fig. 2). In line with these findings, we noticed reduced PK activity and plasma levels in affected patients (Fig. 3b, c). FXII-mediated contact system activation is counter-balanced by C1-INH to prevent continuous kallikrein and bradykinin formation. Thus, our observation of increased FXIIa-C1-INH and PK-C1-INH complexes in FACAS plasma reflected an imbalanced and constitutively active contact system (Fig. 3d, e). Kallikrein cleaves high-molecular-weight kininogen (HMWK) to release bradykinin, a potent but short-lived vasoactive mediator acting on bradykinin B2 receptors. Since bradykinin becomes rapidly degraded by kininases, cleaved HMWK (cHMWK) is used as a surrogate marker of plasma bradykinin formation. Interestingly, we found complete degradation of HMWK in the immunoblot of all tested FACAS patients (Fig. 3f). Correspondingly, the activity of cHMWK was clearly upregulated in patients vs. controls (Fig. 3g). Of interest, bradykinin plasma levels which were available from two FACAS patients revealed higher values compared with healthy controls (Fig. 3h).

To investigate the possibility of local FXII production and activation at sites of inflammation, we evaluated the lesional skin of a FACAS patient following cold exposure. Immunohistochemistry revealed FXII expression by intra- and perivascular neutrophils (but not macrophages; Supplementary Fig. 3) as demonstrated by cytoplasmatic co-staining with antibodies for myeloperoxidase (MPO) and α-FXIIa heavy chain (Fig. 4a). Moreover, we found FXII expression in single peripheral blood neutrophils purified from an affected patient, but not in healthy controls (Fig. 4b). Together, these findings could point toward local FXII production by neutrophils in the skin of FACAS patients. As monocytes and tissue macrophages are known to recruit neutrophils via IL-8 secretion, we hypothesized that contact system mediators might induce the attraction of neutrophils. Accordingly, we found dose-dependent IL-8 release from monocytes treated with increasing doses of bradykinin (Fig. 4c).

FXIIa and bradykinin were previously shown to induce IL-1β secretion from human monocytes and macrophages. Assuming that FXIIa and/or consecutive mediators like bradykinin affect the IL-1β-related phenotype in FACAS, we investigated IL-1β responses in patient samples. Basal IL-1β secretion from patient PBMCs was negligible and comparable with healthy control PBMCs (Supplementary Fig. 4). However, exposure of healthy donor PBMCs to recombinant FXII proteins (wild-type and W268R) suggested an IL-1β priming effect of the FXII W268R mutant protein but not wild-type protein when co-stimulated with the NLRP3 activator ATP (without LPS) (Fig. 5a). To investigate whether the IL-1β priming effect was mediated via FXIIa activity, we assessed the potential of recombinant FXII W268R with an S544A inhibitory mutation (rFXII W268R-S544A) and aFXIIa to induce IL-1β release in PBMCs. Moreover, we blocked protease activity with the small peptide-based inhibitor PPACK (Phe-Pro-Arg-chloro-methylketone). Remarkably, the incapacitated S544A mutant protein showed a similar IL-1β priming effect as the active rFXII W268R (Fig. 5b). In contrast, the priming effect on IL-1β synthesis was absent in PBMCs treated with aFXIIa (Fig. 5c). Also, in LPS-primed and ATP-activated PBMCs, addition of FXIIa or aFXIIa-PPACK did not induce significant changes on IL-1β release (Fig. 5c). Together, these results demonstrate that the priming effect on IL-1β synthesis is not linked to FXIIa activity.

When analyzing biopsies from lesional FACAS skin, we observed pronounced IL-1β expression (Fig. 5d, e). Positive IL-1β staining co-localized with perivascular neutrophils and macrophages, suggesting these cells as the primary source for IL-1β in the skin.

Clinical efficacy of bradykinin B2 receptor antagonism. Based on the collected evidence for excessive contact system activation in FACAS, we initiated a short treatment course over 4 consecutive days with the bradykinin B2 receptor antagonist icatibant in the index patient (patient #1). Individual symptoms were assessed by a patient-reported standardized daily health assessment form that was adapted from a validated tool for CAPS. Within 30 min after the first administration of 30 mg icatibant, the generalized urticarial rash and systemic symptoms improved. The maximum effect (absence of rash and headache, largely improved fatigue) occurred after 90 min and lasted for up to 12 h. Icatibant treatment was initiated during the winter and tolerated well. It completely abrogated skin and systemic symptoms while staying outside for several hours at temperatures below 0 °C. Only arthralgia was still mildly present. Complete relapse of symptoms occurred within 24 h after drug withdrawal (Fig. 6).

Discussion
We report a dominantly inherited cold-induced urticarial auto-inflammatory disease in a four-generation family that is associated with a c.7859A mutation in F12 leading to an amino acid exchange of tryptophan by an arginine (p.W268R). Functional analyses of the mutant protein revealed high susceptibility to
Fig. 2 FXII W268R results in fragmentation and spontaneous activation of FXII. a FXII fragmentation in FACAS plasma samples and FXII W268R mutant proteins. Citrate plasma of affected (#1, #2, #5) and non-affected (#3, #4, #6) family members, recombinant wild-type and W268R FXII expressed in HEK293 cells and plasma from an FXII-HAE patient was separated by SDS-PAGE (reducing blot) and immunoblotted for FXII. Red arrows at 50 kDa indicate heavy chain of cleaved FXII in the plasma of affected patients and recombinant W268R FXII (rFXII W268R), which is barely visible in non-affected and hereditary angioedema (HAE) III plasma or wild-type FXII (rFXII wt). Recombinant W268R and wild-type FXII were expressed in HEK293 cells. Black arrows point to rFXII W268R fragments observed by mass spectrometry in c, n = 5 technical replicates. b Immunoblot (reduced conditions) of recombinant mutant proteins (in the presence or absence of surface activator kaolin) and controls (rFXII wt and purified αFXIIa). In addition to 50 kDa (heavy chain, HC) and 25 kDa (light chain, LC) bands, a further band appears at 60 kDa (II) that is absent in rFXII wt and αFXIIa. Black arrows point to rFXII W268R fragments observed by mass spectrometry in c, n = 3 technical replicates. c Schematic presentation of FXII W268R procession. Red arrows point to mutation at position W268R. Proteolytic sites are indicated by blue arrows. Cleavage after position 353 results in FXIIa W268R HC (50 kDa) and FXIIa W268R LC (25 kDa) fragments. Additional cleavage results in a truncated FXIIa W268R fragment (60 kDa) cleaved at position 447. d FXIIa activity was measured by a chromogenic assay demonstrating considerable spontaneous FXIIa activity in plasma of affected individuals (n = 3), but less in healthy donors (n = 3 healthy family members). e Spontaneous FXIIa activity was also observed in culture supernatants from HEK293 cells expressing W268R, but not wild-type FXII. Graphs show individual data points and the mean ± s.e.m. Statistical significance is indicated by unpaired Student’s t test with Welch’s correction (d, e). Source data are provided as a source data file.
autocatalytic cleavage after residue AA353, which is known to be associated with FXIIa formation. This is paralleled by constitutive activation of the p.W268R variant of FXII resulting in an overactive contact system and release of potent inflammatory cytokines such as IL-1β. The relevance of bradykinin as a critical mediator is underlined by the immediate clinical response to treatment with the bradykinin B2 receptor antagonist icatibant.

Skin and systemic symptoms associated with IL-1β-mediated inflammation in FACAS are reminiscent of the clinical phenotype of CAPS. Its underlying mechanisms, however, are different and share signaling cascades of kallikrein–kinin activation with HAE.

FXII is primarily produced and secreted by hepatocytes. It circulates as a single-chain inactive zymogen in the plasma and converts into FXIIa (composed of an N-terminal heavy chain and a C-terminal light chain) upon contact with negatively charged surfaces such as polyphosphates released by platelets or by limited

Fig. 3 FXII W268R activates the kallikrein–kinin pathway. a Citrate plasma of affected (n = 3) and healthy family members (n = 3) was immunoblated for prekallikrein (PK) showing substantial cleavage of PK (black arrow) to kallikrein red arrow), n = 2 technical replicates. This corresponds to reduced PK activity (b) and plasma levels (c) in affected individuals (n = 3 affected vs. n = 3 healthy). FXIIa-C1-inhibitor (FXIIa-C1-INH) and (e) PK-C1-INH complexes in patient plasma (n = 3 affected vs. n = 3 healthy). Citrate plasma of affected (n = 3) and healthy family members (n = 3) was immunoblotted for high-molecular-weight kininogen (HMWK) showing its complete degradation (black arrow) to cleaved HMWK (cHMWK) (red arrow), n = 2 technical replicates. Correspondingly, increased cHMWK activity (g) and bradykinin plasma levels (h) (n = 3 affected vs. n = 3 healthy; n = 2 affected vs. n = 7 healthy and n = 2 HAE I/II) are detectable in the plasma of affected patients. PK and cHMWK activity were measured by chromogenic assays, PK and bradykinin plasma levels were assessed by ELISA. Graphs show individual data points and the mean ± s.e.m. Statistical significance is indicated by unpaired Student’s t test with Welch’s correction (b, c, d, e, g) or one-way ANOVA with Sidak’s multiple comparisons test (h). Source data are provided as a source data file.
proteolysis\textsuperscript{15}. In the pathogenesis of FXII-HAE, defective glycosylation augments contact-mediated FXII autoactivation, which enhances bradykinin formation via the kallikrein–kinin system\textsuperscript{26,27}. Our data, showing partially cleaved and active FXII in patient plasma as well as enzymatic activity of recombinant W268R in cell culture supernatants, suggest spontaneous, intracellular conversion of FXII into FXIIa in FACAS. In contrast, spontaneous FXII cleavage was absent in FXII-HAE plasma, which is in line with previous reports showing the need for a plasmatic trigger to cleave and activate FXII in HAE\textsuperscript{15}.

The novel mutation induces an exchange of tryptophan to arginine (W268R) inside the kring domain of FXII. Kringles are common in members of the coagulation cascade and are thought to play essential roles in binding target molecules/structures, such as membranes, other proteins, or phospholipids, and in the regulation of proteolytic activity\textsuperscript{26,27}. Our observation of fragmented rFXII-W268R that is cleaved after residue AA353 argues for increased autoactivation and may explain the high spontaneous FXIIa activity observed in plasma and supernatant of HEK293 cells transfected with rFXII-W268R. These findings differ from recent studies in FXII-HAE with p.T309R mutation. This variant was demonstrated to be enzymatically truncated after residue AA309 resulting in the loss of the heavy-chain fragment and dysregulated PK activation\textsuperscript{28}. Our data do not support truncation of W268R after residue AA268. In contrast, we observed additional cleavage after AA447 in the recombinant protein. However, this was not detected in plasma of the affected patients, suggesting that this might be the result of artificial expression in HEK293 cells. From our data, we conclude that high susceptibility to cleavage at the regular cleavage site at AA353/354 leads to continuously increased FXIIa activity. In addition, the exchange of the aromatic and lipophilic tryptophan to the hydrophilic and, at physiological pH, positively charged arginine may foster binding of mutant FXII to negatively charged surfaces, thereby further enhancing its activation state leading to an exhaustion of the contact system with fully cleaved PK and HMWK, and constantly raised bradykinin levels in the patient plasma.

Neutrophil-dominated dermal infiltrates are known from urticarial autoimmune diseases Schnitzler's syndrome and CAPS\textsuperscript{29}. These are classified as neutrophilic urticarial dermatosis as opposed to the rather sparse lympho-histiocytic infiltrates in lesional skin of common urticaria. Their role, however, remains ill-defined. Interestingly, peripheral blood neutrophils were previously shown to produce zymogen FXII, which functionally differs from hepatic FXII and contributes to neutrophil trafficking via cell adhesion, migration, and release of extracellular DNA at sites of inflammation\textsuperscript{30}. Conceivably, the generalized urticarial rash in FACAS could be mediated via neutrophil-derived and pre-activated FXII at the sites of inflammation as suggested by the cytoplasmatic expression of αFXIIa in the skin and peripheral blood neutrophils. IL-8 is known to induce neutrophil recruitment\textsuperscript{19}. Supported by our data, plasma-derived BK may induce IL-8 production by skin macrophages to recruit neutrophils (Fig. 7). We hypothesize that upregulated FXII expression in FACAS neutrophils may be mediated via a paracrine feedback loop involving contact system components such as plasma kallikrein and HMWK. These were both shown to activate (and in
**Fig. 5 FXII W268R induces and increases IL-1β secretion and expression.**

- **a-c** Peripheral blood mononuclear cells were isolated from healthy control subjects by density-gradient centrifugation. Interleukin-1β (IL-1β) secretion requires a two-step process: (i) transcriptional upregulation of pro-IL-1β via toll-like receptors and (ii) conversion of pro-IL-1β into its active form by inflammasome activation. Cells were pre-stimulated with toll-like receptor agonist lipopolysaccharide (LPS), and nucleotide binding like receptor protein 3 (NLRP3) inflammasome activation was induced by the danger molecule ATP (LPS + ATP = blue symbols). Red symbols denote LPS-stimulated cells only, and green symbols represent ATP-activated cells without priming by LPS. Experiments were performed in the presence of FXII-depleted plasma (control) and addition of recombinant FXII wild-type (rFXII wt) and W268R (rFXII W268R) protein (a) or rFXII W268R-S544A (b). Experiments in (c) were conducted with purified FXII, FXIIa, and different doses of the small peptide-based inhibitor PPACK. Black arrows indicate priming effect by the mutant rFXII W268R (a) and the active site incapacitated rFXII W268R-S544A (b), which was absent in rFXII wt and control. Also, the presence of FXIIa (with or without PPACK) did not induce changes in IL-1β release (c). IL-1β secretion was assessed by ELISA. Pooled data of \( n = 3 \) different (a) or \( n = 2 \) (b, c) experiments. Bar graphs show individual data points and the mean ± s.e.m. Statistical significance in is indicated by one-way ANOVA (a, c) with Tukey’s multiple comparisons test for ATP between the groups control vs. rFXII W268R and rFXII wt vs. rFXII W268R (a) or unpaired Student’s t test (b). d, e Representative image from serial sections of the lesional skin from FACAS patient following cold exposure exhibits IL-1β immunoreactivity. IL-1β expression is pronounced around blood vessels (yellow arrows) and co-localizes with neutrophils (d) and macrophages (e) (white arrows). Original magnification ×400, \( n = 2 \) technical replicates. Source data are provided as a source data file.
case of HMWK also bind to) neutrophils\textsuperscript{31,32}. Thus, the high degree of contact system activation could be the reason why FXII expression was restricted to FACAS and not observed in healthy control neutrophils.

The sensitivity to cold as a trigger and amplifier of clinical symptoms in FACAS is known from autoinflammatory diseases such as CAPS. Here, cold was proposed to escalate constitutive inflammasome activity via increased production of reactive oxygen species induced by cellular stress to temperature shifts\textsuperscript{33}. In FACAS, however, inflammasomes of mononuclear cells are not constitutively activated (Supplementary Fig. 4), suggesting alternative mechanisms. Of interest, C1-INH activity was previously reported to be inhibited by cold as shown by complete plasma HMWK cleavage at 0 °C in vitro\textsuperscript{34}. We assume that cold-induced decreases in tissue C1-INH activity may contribute to an imbalanced contact system in FACAS. Temperatures above a certain threshold (clinically around 20 °C) may protect from inflammatory symptoms because of adequate local C1-INH activity that counterbalances kallikrein activation. Based on this hypothesis, lower temperatures will reduce C1-INH activity leading to spontaneous but locally restricted activation of the kallikrein-kinin pathway and bradykinin production in the tissue, sufficient for wheal induction but not angioedema.

Bradykinin drives the inflammation in experimental models of osteoarthritis and neuroinflammation via upregulation of cytokines such as IL-1\textbeta\textsuperscript{35,36}. Vice versa, IL-1\textbeta is known to augment the accumulation of bradykinin via upregulation of bradykinin receptors\textsuperscript{37,38} and activation of prekallikrein–HMWK complexes\textsuperscript{39}. Also, IL-1\textbeta downregulates angiotensin-converting enzyme (ACE) expression in endothelial cells, resulting in less efficient bradykinin elimination\textsuperscript{40}. Accordingly, elevated skin and systemic IL-1\textbeta levels in FACAS may amplify inflammatory responses via a positive feedback loop that induces further bradykinin production. Moreover, bradykinin was shown to activate

**Fig. 6** The bradykinin B2 receptor antagonist icatibant resolves FACAS signs and symptoms. During winter with continuous skin symptoms, the index patient administered icatibant 30 mg. a Urticarial rash completely resolved within 90 min post icatibant application (from the left to right). b Total and individual symptoms reported and assessed daily over a time period of 8 days by the patient using a standardized daily health assessment form covering the main symptoms rash, fatigue, headache, arthralgia, and fever/chills. The total score ranges from 0 = no symptoms to 50 = maximum of symptoms. Each subscore can range from 0 = no symptoms to 10 = maximum of symptoms. The subscores do not show fever/chills as these symptoms were not present at that time. Icatibant was given on 4 consecutive days (red arrows).
and degranulate mast cells in vitro resulting in the release of heparin and polyphosphates, which may facilitate the local production of bradykinin via FXII activation. Also, skin mast cells contribute to IL-1β-mediated inflammation in urticarial autoimmune inflammatory diseases, and their abundant expression and activation may promote the urticarial phenotype in FACAS patients (Fig. 7).

The clinical response to treatment with the IL-1 receptor antagonist anakinra mirrors the increased presence of IL-1β in FACAS tissue similar to other urticarial autoimmune inflammatory diseases such as CAPS. Incomplete clinical improvement by anakinra may be explained by continuous bradykinin generation via complex feedback loops and possibly further cytokines such as TNFα, which are induced by bradykinin and not inhibited by IL-1 blockade. Of further interest, our findings suggest that the p.W268R variant can prime cells for IL-1β production via pattern-recognition receptors, without the need of an exogenous stimulus by pathogen-associated molecular patterns such as LPS. Surprisingly, the IL-1β priming effect is not linked to FXIIa activity as shown by the missing impact of αFXIIa on IL-1β secretion. Thus, we believe that conformational changes of FXII p.W268R result in the recognition as a danger signal that serves as toll-like receptor agonist and inducer of IL-1β release.

The prompt and striking clinical response to the bradykinin B2 receptor antagonist icatibant is in line with its efficacy in HAE patients and strongly supports our model of a bradykinin-mediated mechanism. Reasons for the persistence of mild arthralgia may be the short treatment duration of icatibant or the fact that bradykinin effects on joint inflammation are mediated via the B1 receptor rather than the B2.

Limitations of our study include the small patient and sample sizes. Until now, there are no other families or individuals known with FACAS. Hence, it will be important to screen patients with similar phenotypes for the FXII p.W268R mutation in order to confirm our data. The mechanism by which cold induces inflammation as well as the role of neutrophils in FACAS need to be further evaluated. Also, the duration of treatment with anakinra and icatibant was only short due to side effects and limited efficacy (anakinra) or limited availability (icatibant). Data on long-term efficacy are still missing.

In conclusion, the pathomechanisms underlying this novel disease illustrate the critical role of fragmented and pre-activated FXII in driving skin and systemic inflammation as well as the close link between contact system activation and cytokine signaling. Therapeutic approaches in FACAS should aim at continuous control of inflammation in order to prevent long-term sequelae such as amyloidosis. Based on the short half-life of icatibant of few hours, inhibitors of contact activation including kallikrein antagonists and FXII antibodies are promising drug candidates of interest.

Methods

Patients. A total of eight family members, four of them affected and four non-affected, were analyzed. All of them provided written informed consent. This included the publication of individual data. For affected children, parents provided written informed consent, respectively. The authors affirm that human research participants provided informed consent for publication of the images in Figs. 1a, 6a. The study was approved by the local Charité Ethics Committee, Charité—Universitätsmedizin Berlin, Germany.

Genetic studies. Genomic DNA was isolated from saliva samples of eight family members, and whole-exome sequencing (SureSelect human all exon v5.0, Agilent, Santa Clara, California) of three affected family members across three generations was performed and analyzed as described before in greater detail. Briefly, genetic variants in the candidate genes were filtered to be present in all three sequenced individuals (in any zygosity) and to occur at ≤5% in 1000 Genomes database. DNA was available for follow-up Sanger sequencing from seven family members including all four affected individuals to determine consistency of variant segregation with disease. DNA exome capture was performed using the SureSelect human all exon v5 kit according to the manufacturer’s protocol and the library sequenced on the HiSeq 2000 platform (Illumina, San Diego, California), generating data sufficient to provide ≥50× coverage across the exome. Raw data were aligned using Novoalign v2.08 and duplicate reads marked using Picard v1.97. Variant calling was performed using Samtools v0.1.19 and variants annotated using Annovar (23rd August 2013 release) utilizing ReSeq gene definitions. Candidate genes were identified in HGMD release 2015.4 using the search term urticaria.
Table 2 Primer sequences for FXII genetic constructs.

| N-Frag_For | GGAAGATGCAAACCTGTTATTTGCTGATTTCCACG |
| N-Frag_Rev | GCTCCGGGGTGCTGGAGCTAGGGGGG |
| M-Frag_For | GACCCGGGACGCCCTGGCCGACGAG |
| M-Frag_Rev | CGTGTTGCTGGAGGTCGGCCGCGGGCGTGG |
| C-Frag_For | GACCTCGGAGACCACCTGGCTGCAGG |
| C-Frag_Rev | CGAGCCGGGCGCTCATGCAAGAAGCTGTCCTCG |

Separation analysis was performed on amplicons generated using primers forward S′-CTAACCTCCGGGCTGC-3′ and reverse 5′-CGACCCGGGAGGTTAG-3′ using standard techniques. Unidirectional Sanger sequencing was performed using the forward primer.

Blood collection. Following standard procedures, nine volumes of blood were collected in one volume of sodium citrate. (3.2% w/v). Samples were centrifuged twice at 2000g for 10 min. Plasma from FACAS-affected family members and from healthy donors was stored in aliquots at −80 °C until use. For PBMC stimulation assays, blood from FACAS-affected family members or healthy control donors was sampled into lithium heparin tubes and processed as described above. Citrate and heparin tubes were from Greiner Bio One (Kremsmünster, Austria). For plasma bradykinin level assessment, blood was sampled into SCAT-169-4.5/5 protase inhibitor blood collection tubes (Haematologic Technologies, Essex Junction, Vermont), centrifuged at 2000 × g for 10 min and stored as described above.

Western blot. Frozen plasma of affected and healthy individuals was thawed on ice, diluted 1:10 (FXII), 1:20 (HMWK), or 1:40 (PK) in sample buffer (15.5% glycerol, 96.8 mM Tris-HCl, 3.1% SDS, and 0.003% bromophenol blue, 25 µM DTT (Sigma-Aldrich, St. Louis, Missouri)), boiled for 5–10 min and briefly chilled on ice. Recombinant constructs were treated similarly. Approximately 50–100 ng protein of recombinant proteins or 5 µl protein sample was loaded and separated on 10% Tris/glycine or Bolt 4–12% Bis-Tris Plus Gels (Merck-Millipore) and separated at 200 V (running buffer 190 mM glycine, 25 mM Tris, 0.1% SDS) or 165 V (MOPS running buffer (Merck-Millipore, Amsterdam, The Netherlands)), respectively. Bands were visualized with 0.1% CoaChrom Diagnostica) as specified by the manufacturer instructions and destained using 10% acetic acid, 20% ethanol. Dehydrated bands were cut by scalpel, and sent for mass spectrometry analysis using an EasyLC 1200 system (Thermo Fisher Scientific) on in-house manufactured 20-cm cisliss silica microcolumns packed with ReproSil C18 resin (inner diameter 75 µm, Dr. Maisch GmbH, Ammerbuch-Entringen, Germany).

Peptides were separated using a 40% acetonitrile gradient (20 µl eluate) at a nanoflow rate of 250 nl/min. Eluting peptides were directly ionized by electrospray ionization and analyzed on a Q Exactive HF-X (Thermo Fisher Scientific). Mass spectrometry was performed in data-dependent positive mode with one full scan (m/z range ~350–1800; R = 60000, target value: 3 × 106; maximum injection time = 45 ms). The Top 20 most intense ions with a charge state of 1 and 2 were selected (R = 45000, target value = 1 × 106; isolation window = 1.3 m/z; maximum injection time = 86 ms). Dynamic exclusion was performed for selected precursor ions was set to 20 s. Data analysis was performed using MaxQuant software package (version 1.8.0.8). The internal 40–45% search engine was used to search MS2 spectra against human coagulation factor XII wild-type and W268R mutant sequence containing forward and reverse sequences. A database for contaminants was included. The search included variable modifications of methionine oxidation, N-terminal acetylation, deamidation (N and Q), and the fixed modification of carbamidomethyl cysteine. Minimal peptide length was set to six amino acids, digestion parameter was semitryptic. The FDR was set to 0.01 for peptide, protein, and site identifications. To filter for confidently identified peptides, a minimum of two MS counts and valid value in both replicates for at least one sample group were required. Intensity values were log2 transformed. Mass spectrometry was performed twice; n = 2 duplicates were assessed for each band.

Factor XII protein expression and purification. The eukaryotic expression vector pDNA6/V5-His A (Thermo Fisher Scientific) was modified by insertion of a murine Igk secretion signal and two strep-tags through the HindIII and EcoRI digestion sites of the cysteine residues (called pSM2). cDNA of the F12 gene was kindly donated by Dr F. Citarella and matches the National Center for Biotechnology Information reference sequence (NM_000505.3), except for the silent mutation c.tIl1104T > C (P460P, mature protein numbering). A sequence coding for a tobacco etch virus (TEV) protease cleavage site was inserted from the F12 sequence by using PCR with the Primers TEVP (Thermo Fisher Scientific). FXII WT cDNA was digested through two-step assembly of three double-stranded DNA segments (annotated as N, M, and C, respectively; Primer sequences in Table 2). Fragment N with the desired mutation (T859A) was ordered as double-stranded DNA fragments from Integrated DNA Technologies (Coralville, Iowa) and ligated into the pET2.1/blunt vector (Thermo Fisher Scientific) according to the manufacturer instructions and amplified through PCR. Fragments M and C were amplified by PCR from FXII-WT cDNA in pSM2. First, the N and M segments were digested with Xmal and subsequently, fused by ligation with T4 ligase (Thermo Fisher Scientific). The fused product (NM) was amplified via PCR. Hereafter, the NM and C segments were digested with Bsal and fused together by ligation with T4 ligase. The assembled cDNA product was ligated into the pET2.1/blunt vector according to the manufacturer’s instructions. Inserts were digested with EcoRI and NotI and ligated into the EcoRI and NotI-restricted pSM2 vector. All restricted clones were purchased from New England Biolabs (Ipswich, MA). Sequences were verified by Sanger sequencing (Macrogen Europe, Amsterdam) with a charge state between +1 and +2. HEK293F (LGC Standards GmbH, Wesel, Germany) cells were transfected using 293Fectin (Thermo Fisher Scientific) and cultured in Gibco Freestyle 293 Expression Medium (Thermo Fisher Scientific) at a concentration of 1 × 10^6 cells/ml in 2 ml. Four days after transfection, the cell suspension was centrifuged and 500 µg of the supernatant was collected and stored until use at −20 °C. For protein purification, transfection was repeated in a volume of 1× cell suspension, and culture medium was harvested after 1 week. FXII-W626R was purified via a strep-tag using Strep-Tactin Sepharose beads (IBA-Life Sciences, Goettingen, Germany) and dialyzed overnight at 4 °C against 4 mM sodium acetate–HCl with 150 mM NaCl (pH 5.5) using a Spectra/Por dialysis membrane (molecular weight cutoff 3.5 kDa, Spectrum Labs, Rancho Dominguez, California) which was preincubated with polybrene (Sigma-Aldrich).

Chromogenic assays. Chromogenic substrate H-D-Pro-Phe-Arg-pNA (0.5 mM, Bachem, Bubendorf, Switzerland) was used to detect FXII activity from recombinant constructs in culture medium. Ninety-six-well Costar plates (Corning, New York) were blocked with HBS (10 mM HEPES (VWR International, Amsterdam, The Netherlands), 150 mM NaCl, 1 mM MgSO4, 5 mM KCl, pH = 7.4) containing 0.5% BSA. Subsequently, 30 µl of culture medium supernatant containing rFXII was added followed by 10 µl of substrate. Substrate conversion was immediately read out at 405 nm at 37 °C. FXII activity in plasma was assessed using the Chromogenic Assay for Factor XIIa-like Activity (COA0088, CoaChrom Diagnostica, Maria Enzersdorf, Austria) as specified by the manufacturer. Absorbance signals were measured in the VIKTOR X5 (PerkinElmer, Waltham, MA), and analyzed using GraphPad Prism 7 or Microsoft Excel 2016.
PK activity (routine lab) and PK ELISA. Prekallikrein activity was assessed by coagulometric assay in routine assays at Labor Berlin (1335 Berlin, Germany). Prekallikrein concentration was measured using the Prekallikrein and Kallikrein Human ELISA (ab171015, Abcam, Cambridge, UK) as specified by the manufacturer. Absorbance signals were measured in the VIKTOR X5 (PerkinElmer, Waltham, MA), and analyzed using GraphPad Prism 7.02 or Microsoft Excel 2016.

Enzyme-C1-INH complexes ELISA. Methods on V1,H (variable domain heavy-chain antibody–dye conjugate) detection and production, and execution for this assay are described earlier by de Maat25. In brief, V1,H 1B12 (5 µg/mL), a monoclonal V1,H directed against complexed C1-INH, was immobilized on 96-well MaxiSorp plates (Corning). Plasma samples were diluted in 10 mM HEPES, 150 mM NaCl, 1 mM MgSO4, 5 mM KCl, pH = 7.4 (HBS) supplemented with 0.1% (w/v) Tween-20 and 1% (w/v) skimmed milk powder (Sigma-Aldrich) and containing 200 µM F-Pro-Arg-chloromethylketone (PPACK, HaematoLogic Technologies). For reference, normal pooled plasma from healthy donors (NPP) activated with DXX (average molecular weight 500,000, Sigma-Aldrich) for 30 min as well as non-activated NPP were prepared in HBS containing PPACK and mixed to generate samples containing 0–100% DXX-activated NPP. Samples were incubated on the plate for 1 h. Enzyme present in the captured complexes was detected with a biotinylated monoclonal V1,H H targeted against the jFXXIA (5 µg/mL, made and biotinylated in-house36) followed by streptavidin-poly-HP-R 1:4000 (M2051, Sanquin Blood Supply, Amsterdam, The Netherlands) or a polyclonal sheep anti-PIK IgG 1:2000 (SAPK-AP, Afshouse47) followed by streptavidin-poly-HRP 1:4000 (M2051, Sanquin Blood Supply, Amsterdam, The Netherlands) or a polyclonal VHH targeted against the enzyme-C1-INH complex antibodies. Absorbance signals were measured in the VIKTOR X5 and analyzed using GraphPad Prism 7.02 or Microsoft Excel 2016 against Bradykinin standard curves.

Cleaved kinogen activity ELISA. Methods on V1,H selection, production, and execution for this assay are described more detailed by Hofman et al.48. In brief, patients samples were diluted 64-fold in DPBS (Thermo Fisher Scientific) supplemented with 0.1% (w/v) Tween-20 and 1% (w/v) skimmed milk powder and containing 200 µM PPACK and 0.5 µg/ml DXS to generate HMWK complexes. For reference, HMWK complexes were prepared in HBS containing PPACK and mixed to generate samples containing 0–100% HMWK-activated NPP. Samples were incubated on the plate for 1 h. Enzyme present in the captured complexes was detected with a biotinylated monoclonal V1,H H targeted against the jFXXIA (5 µg/mL, made and biotinylated in-house36) followed by streptavidin-poly-HP-R 1:4000 (M2051, Sanquin Blood Supply, Amsterdam, The Netherlands) or a polyclonal VHH targeted against the enzyme-C1-INH complex antibodies. Absorbance signals were measured in the VIKTOR X5 and analyzed using GraphPad Prism 7.02 or Microsoft Excel 2016 against Bradykinin standard curves.

Bradynkin ELISA. Bradykinin levels in patient and healthy control plasma were assessed by ELISA (ADI-900-206, Enzo Life Sciences, Farmingdale, New York) as described by the manufacturer. Absorption signals were measured in the VIKTOR X5 and analyzed using GraphPad Prism 7.02 or Microsoft Excel 2016 against Bradykinin standard curves.

IL-1β and IL-8 ELISA. IL-1β and IL-8 secretion was assessed by ELISA (Duoset, DY201/DY208-05, R&D Systems, Minneapolis, MN) as specified by the manufacturer. For greater precision, ELISAs were carried out at ¼ of the recommended volumes per well in black NUNC 384-well MaxiSorp plates (Thermo Fisher Scientific) for 1 h. Enzyme present in the captured complexes was detected with a biotinylated monoclonal anti-CD163 (K5005, Dako) for MPO staining or labeled polymer-HP-R anti-rabbit (K4011, Dako) together with AEC + high sensitivity substrate chromogen readout and antibody served as negative controls. Sections were analyzed by fluorescence microscopy using a Zeiss Axioplan 2 with Axiosvision software 4.8.1. Images of cytopsins were further analyzed by ImageJ 1.52a.

PBMC stimulation assays. Venous blood from patients and healthy donors was sampled into heparin tubes. In all, 15 ml of Ficoll-Paque solution (GE Healthcare) were pipetted into 50-ml Leucosep tubes (Greiner Bio One) and centrifuged at 1000 g for 4 min. Up to 20 ml of blood sample per tube was added, filled up to 50 ml with DPBS and centrifuged for 10 min at 1000 x g with 33% acceleration and without breaks. The upper plasma layer was discarded, the PBMC layer was collected and filled into a new 50-ml tube. PBMCs were washed twice by adding ad 50 ml DPBS, centrifugation for 10 min at 350 x g, and resuspension in residual DPBS. After the washing steps, 10 ml of PBMC medium (RPMI 1640 containing 1% FBS (both from Biochrom, Berlin, Germany), 100 U/ml penicillin, 100 µg/ml streptomycin (HyClone 100X Pen/Strep, GE Healthcare)) was added, and cells were counted using trypan blue staining and hemocytometer. PBMC number was adjusted to 3 x 10^6/ml, and 100 µl of the suspension were added per well to Falcon TC96-well plates (Corning). After 2 h incubation at 37 °C and 5% CO2, the supernatant containing non-monoocytes was discarded, and to the adherent monocytes, 100 µl of containing 0.5 µg/ml LPS (E. coli R515, Enzo Life Sciences) or not was added for NF-κB priming. Monocytes were incubated for 20 h at 37 °C and 5% CO2, supernatant was discarded, and 20 µl per well of PBMC medium was added. In all, 20 µl of either bradykinin at stated final concentrations or FXII constructs (with or without stated final concentrations of PPACK) diluted in PBMC medium were added at a final concentration of 35 ng/µl, followed by incubation at 37 °C and 5% CO2 for 1 h. Subsequently, 10 µl of 10 mM ATP (Sigma Aldrich) diluted in PBMC medium were added to a final concentration of 2 mM for inammmasome activation, followed by another 1 h incubation at 37 °C and 5% CO2. Supernatants were collected and stored at –20 °C until analysis by ELISA.

Statistical analysis. Sample sizes were limited due to patient availability. The number of samples (all assessed in duplicates) and experiments are displayed and listed in the figure legends. The use of statistical tests is indicated in the figure legends as well.

Reproting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. Source data for figures 2a–e, 3a–c, 4b, 6c, 5a–c, and Supplementary Figs. 1, 2 and 4 are provided as a Source Data file (https://doi.org/10.6084/m9.figshare.13117669). The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE [1] partner repository with the dataset identifier PXD016341. As the informed consent obtained from FACAS patients does not allow for public deposition of the sequencing data, the filtered WES sequencing data from patients #2, #5, and #7 can be communicated upon reasonable request to R.J.P. and S.E. All other relevant data are contained within the article and its Supplementary Information.

Received: 13 March 2019; Accepted: 5 December 2019; Published online: 10 January 2020.
References

1. Hoffman, H. M., Mueller, J. L., Broide, D. H., Wanderer, A. A. & Kolodner, R. D. Mutation of a new gene encoding a putative pyrin-like protein causes familial cold autoinflammatory syndrome and Muckle-Wells syndrome. *Nat. Genet.* **29**, 301–305 (2001).

2. Toplak, N. et al. An international registry on autoinflammatory diseases: the Eurofever experience. *Ann. Rheum. Dis.* **71**, 1177–1182 (2012).

3. Naka, M. et al. Somatic NLRP3 mosaicism in Muckle-Wells syndrome. A genetic mechanism shared by different phenotypes of cryopyrin-associated periodic syndromes. *Ann. Rheum. Dis.* **74**, 603–610 (2015).

4. Tanaka, N. et al. High incidence of NLRP3 somatic mosaicism in patients with chronic infantile neurologic, cutaneous, arthritic syndrome: results of an International Multicenter Collaborative Study. *Arthritis Rheum.* **63**, 3265–3262 (2011).

5. Jeru, I. et al. Mutations in NALP12 cause hereditary periodic fever syndromes. *Nat. Genet.* **46**, 1140–1146 (2014).

6. Romberg, N. et al. Mutation of NLRCA causes a syndrome of enteroctolitis and autoinflammation. *Nat. Genet.* **46**, 1135–1139 (2014).

7. Ormstron, N. et al. Clinical presentation and pathogenesis of cold-induced autoinflammatory disease in a family with recurrence of an NLRP12 mutation. *Arthritis Rheum.* **63**, 830–839 (2011).

8. Ombretin, M. J. et al. Cold urticaria, immunodeficiency, and autoimmunity related to PLCG2 deletions. *N. Engl. J. Med.* **366**, 330–338 (2012).

9. Stenson, P. D. et al. The Human Gene Mutation Database: towards a comprehensive repository of inherited mutation data for medical research, genetic diagnosis and next-generation sequencing studies. *Hum. Genet.* **136**, 665–677 (2017).

10. Løk, M. et al. Analysis of protein-coding genetic variation in 60,706 humans. *Nature* **536**, 285–291 (2016).

11. Cichon, S. et al. Increased activity of coagulation factor XII (Hageman factor) causes hereditary angioedema type III. *Am. J. Hum. Genet.* **79**, 1098–1104 (2006).

12. Verene, C. L. et al. Hereditary angioedema with normal C1 inhibitor and F12 mutations in 42 Brazilian families. *J. Allergy Clin. Immunol.* **6**, 1209–1216 e1208 (2018).

13. Schoeless, M. et al. Mutations in the human factor XII gene. *Blood* **90**, 3967–3977 (1997).

14. Renne, T., Schmaier, A. H., Nickel, K. F., Blomback, M. & Maas, C. In vivo contact activation: effects of surface and inhibitors. *Med. Prog. Technol.* **15**, 53–62 (1988).

15. Xie, Z., Dai, J., Yang, A. & Wu, Y. A role for bradykinin in the development of anti-collagen antibody-induced arthritis. *Rheumatology* **53**, 1301–1306 (2014).

16. Niewiarowska-Sendo, A., Kozik, A. & Guevara-Lora, I. Kinin peptides enhance inflammatory and oxidative responses promoting apoptosis in a Parkinson’s disease cellular model. *Mediators. Inflamm.* **2016**, 4567343 (2016).

17. Passos, G. F. et al. Kinin B1 receptor up-regulation after lipopolysaccharide administration: role of proinflammatory cytokines and neutrophil influx. *J. Immunol.** **172**, 1839–1847 (2004).

18. Rehbock, J., Miska, K. & Buchinger, P. Induction of the bradykinin B2-receptor, but not of the bradykinin B1-receptor, by interleukin-1 beta in cultivated human decidua-derived cells. *Immunopharmacology* **43**, 235–239 (1999).

19. Joseph, K., Tholanikunnel, B. G. & Kaplan, A. P. Cytokine and estrogen stimulation of endothelial cells augments activation of the prekallikrein-high molecular weight kininogen complex: Implications for hereditary angioedema. *J. Allergy Clin. Immunol.** **108**, 375–381 (2001).

20. Sajzoffma, O., Nyman, T. & Fhyurquist, F. Downregulation of angiotensin-converting enzyme by tumor necrosis factor-alpha and interleukin-1 beta in cultured human endothelial cells. *J. Vasc. Res.** **38**, 370–378 (2001).

21. Oschatz, C. et al. Mast cells increase vascular permeability by heparin-initiated bradykinin formation in vivo. *Immunity* **34**, 258–268 (2011).

22. Moreno-Sanchez, D., Hernandez-Ruiz, L., Ruiz, F. A. & Docampo, R. Polyphosphatase is a novel pro-inflammatory regulator of mast cells and is located in acidocalcisomes. *J. Biol. Chem.** **287**, 28435–28444 (2012).

23. Nakamura, Y. et al. Mast cells mediate neutrophil recruitment and vascular leakage through the NLRP3 inflammasome in histamine-independent urticaria. *J. Exp. Med.** **206**, 1037–1046 (2009).

24. Sundqvist, G. & Lerner, U. H. Bradykinin and thrombin synergistically potentiate interleukin 1 and tumour necrosis factor induced prostanoid biosynthesis in human dental pulp fibroblasts. *Cytokine** **8**, 168–177 (1996).

25. Pengelly, R. J. et al. Resolving clinical diagnoses for syndromic cleft lip and/or palate phenotypes using whole-exome sequencing. *Clin. Genet.** **88**, 441–449 (2015).

26. Cox, J. & Mann, M. MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. *Nat. Biotechnol.** **26**, 1367–1372 (2008).

27. de Maat, S., van Doornemen, S., de Groot, P. G. & Maas, C. A nonobesity-based method for tracking factor XII activation in plasma. *Thromb. Haemost.* **104**, 458–468 (2013).

28. Hofman, Z. L. M. et al. Cleaved kininogen as a biomarker for bradykinin release in hereditary angioedema. *J. Allergy Clin. Immunol.** **140**, 1700–1703 e1708 (2017).

29. Scheffel, J. M. et al. Cold-induced urtiaral autoinflammatory syndrome related to factor XII activation. [https://doi.org/10.6084/m9.684421](https://doi.org/10.6084/m9.684421)

Acknowledgements

This work was supported by the Fritz Thysen Foundation (to K.K.). M.M. acknowledges support by the DFG (MA1919/14-1). C.M. acknowledges support from the Landsteiner Foundation for Innovative Blood Transfusion Research and the Netherlands Thrombosis Foundation, Z.L.H. is supported by the Alexandre Suerman Programme of the Utrecht University Medical Center. We thank Evelyn Hagen for excellent technical assistance. Most of all, we would like to thank the family members for their great support and cooperation.

Author contributions

J.S.: supervision of experiments, critical input on study concept and paper. N.M.: performed cell stimulation assays on IL-1β. Z.L.H. is supported by the Alexandre Suerman Programme of the Utrecht University Medical Center. Support by the DFG (MA1919/14-1). C.M. acknowledges support from the Landsteiner Foundation for Innovative Blood Transfusion Research and the Netherlands Thrombosis Foundation. Z.L.H. is supported by the Alexandre Suerman Programme of the Utrecht University Medical Center. We thank Evelyn Hagen for excellent technical assistance. Most of all, we would like to thank the family members for their great support and cooperation.

[https://doi.org/10.6084/m9.684421](https://doi.org/10.6084/m9.684421)
analysis and analysis of sequencing data. J.H.: critical input on concept for genetic analysis M.K.: performed and analyzed mass spectrometry data. P.M.: supervised and discussed mass spectrometry analysis. M.K.C.: critical input on manuscript. M.Mau.: critical input on study concept and paper. C.M.: supervision of experiments on contact system and critical input on study concept. K.K.: overall study concept and idea; supervision of experiments and collection of clinical data; data analysis; wrote the paper with contributions from all authors. All authors critically reviewed the paper and approved the final version before submission.

Competing interests
M.Mau. received consultation fees and research support from Shire/Takeda outside the submitted work. K.K. received research support from Shire/Takeda outside the submitted work. The remaining authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41467-019-13984-8.

Correspondence and requests for materials should be addressed to M.M.

Peer review information Nature Communications thanks the anonymous reviewers for their contribution to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.