Hyper-IgE and Carcinoma in CADINS Disease

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Background: Atopic dermatitis (AD) affects up to 25% of children and 10% of adults in Western countries. When severe or recurrent infections and exceedingly elevated serum IgE levels occur in AD patients, an inborn error of immunity (IEI) may be suspected. The International Union of Immunological Societies classification lists variants in different genes responsible for so-called Hyper-IgE syndromes. Diagnosing an underlying IEI may influence treatment strategies.

Methods: Clinical and diagnostic workup of family members are presented including a detailed immunological description and histology of the carcinoma. Functional testing of the novel variant in CARD11 underlying ‘CARD11-associated atopy with dominant interference of NF-kB signaling’ (CADINS) was performed.

Results: We report on an 18-year-old patient with a long-standing history of infections, accompanied by hypogammaglobulinemia, intermittent agranulocytosis, atopy, eosinophilia and colitis. The working diagnosis of common variable immunity was revised when a novel heterozygous CARD11 variant [c.223C>T; p.(Arg75Trp)] was identified. Functional studies confirmed this variant to have a dominant negative (DN) effect, as previously described in patients with CADINS. Five other family members were affected by severe atopy associated with the above variant, but not hypogammaglobulinemia. Malignancies occurred in two generations: an HPV-positive squamous cell carcinoma and a cutaneous T-cell lymphoma. So far, one patient is under treatment with dupilumab, which has shown marked benefit in controlling severe eczema.

Conclusion: The phenotypic spectrum associated with heterozygous CARD11 DN mutations is broad. Partial T-cell deficiency, diminished IFN-γ cytokine and increased IL-4 production, were identified as disease-causing mechanisms. Malignant disease
INTRODUCTION

Atopic dermatitis (AD) is one of the most common childhood diagnoses in Western European countries. It affects up to 25% of children and 10% of adults (2). AD is a common type of eczema; other atopic diseases (e.g. asthma, rhinitis, food allergies) are often associated. Certain inborn errors of immunity (IEI) may also present with eczema and highly elevated serum IgE levels. However, patients with classic IEI such as severe combined immunodeficiency (SCID) or with Omenn, Wiskott-Aldrich, DiGeorge syndromes or DOCK8 deficiency usually manifest in early infancy or childhood, accompanied by infection susceptibility in addition to early-onset severe AD. Other so-called Hyper IgE syndromes (HIES) include susceptibility to bacterial, fungal or viral infections such as molluscum contagiosum and the human papilloma virus (HPV). These include classical autosomal dominant HIES due to STAT3 loss-of-function (LOF) and autosomal recessive entities linked to variants in ZNF341, PGM3, and genes more recently associated with HIES such as SOCS1 genes and CARD11 (Caspase recruitment domain family member 11) (3–7). Autosomal dominant CARD11 variants were first published in 2017 in children and teenagers with the triad of AD/asthma, Hyper-IgE and recurrent infections (8).

CARD11, a multi-domain scaffold protein, belongs to the membrane-associated guanylate kinase (MAGUK) protein family as well as the caspase-associated recruitment domain (CARD) protein family. The protein structure is illustrated in Supplementary Figure 1. As a constituent of the CARD11-BCL10-MALT1 (CBM) complex, it plays a critical role in intracellular signaling cascades, most notably NF-κB activation downstream of antigen receptor signalling in lymphocytes. The phosphorylation of CARD11 leads to a conformational change which allows CARD11 to recruit cofactors including BCL10, MALT1 and TRAF6. This multi-protein signaling complex then activates the IKK complex, leading to phosphorylation of the inhibitory IκBα protein, its ubiquitinylination, and subsequent proteasomal degradation. Activated NF-κB translocates to the nucleus and regulates transcription of proliferative, pro-inflammatory, and anti-apoptotic genes essential for lymphocyte function. The CBM complex is also involved in the activation of c-Jun N-terminal kinase (JNK) and mechanistic target of rapamycin (mTORC1) signalling following T cell receptor (TCR) engagement (9).

The phenotypic spectrum for patients with germline CARD11 DN variants has only been reported sporadically. HPV vaccination in teenage years, and cytology screening analogous with routine cervical swabs may be recommended. Treatment with dupilumab, a monoclonal antibody blocking interleukin-4- and interleukin-13 signaling, may be of benefit in controlling severe and extended AD for some patients as reported for STAT3 loss-of-function.

Keywords: CARD11 deficiency, severe eczema, dupilumab, HPV driven carcinoma, anal carcinoma, mycosis fungoides, hyper-IgE-syndrome, CADINS

RESULTS

Patient Clinical and Immunological Characteristics

Here we describe a novel CARD11 variant (p.Arg75Trp) segregating with Hyper-IgE associated atopy, infections and malignancy in a three-generation kindred with at least 6 affected family members (Figures 1A, B).

The now 18-year-old-index patient (III.4) initially presented at 2 months of age with mastoiditis, pneumonia and agranulocytosis. It was supposed to be caused by allo- or autoantibodies as neutrophil counts normalized after one year. At 2 years of age, he showed a rectal herpesvirus infection. Beginning at age 3, he suffered from recurrent respiratory infections and bronchial obstruction, and was therefore diagnosed with asthma. Severe eczema developed on his hands and feet (Figures 2A, B), as well as recurrent nail mycoses. He was additionally diagnosed with chronic polyposis, sinusitis and food allergies. From 12 years of age onwards, the patient complained of gastrointestinal symptoms, such as abdominal pain and nausea associated with weight loss.

At age 5, hypogammaglobulinemia was diagnosed, and the patient was started on regular subcutaneous immunoglobulin substitution. Although his susceptibility to infection markedly decreased, bronchiectasis was documented at age 13 years. Following several episodes of mild hematochezia, eosinophilic colitis was detected by endoscopy at age 16.
The maternal grandfather (I.1) had suffered from pronounced AD and Mycosis Fungoides (MF) with extracutaneous dissemination, and died at age 56.

The index patient’s mother (II.2) suffered from severe hand and foot eczema, and recurrent respiratory infections. At age 45, she was diagnosed with anal squamous cell carcinoma induced
by a local HPV18 infection (Figure 3). This carcinoma was discovered only after it had already metastasized, and despite removal of all intra-pelvic organs, three relapses occurred within 5 years. Clinical management was switched to palliative care.

The index patient’s maternal aunt (II.1) suffered from warts as well as from hand and foot eczema, allergies and asthma from her teenage years. Her son (III.1) and daughter (III.2) – the index patient’s cousins - report various food allergies, bronchial asthma and severe AD (Figures 2C, D). One of them (III.1) was successfully treated with the humanized anti-IL4Ra biological drug dupilumab, which blocks both IL-4 and IL-13 signaling. Photodocumentation pre- and post-treatment is presented in Figures 2E, F. The cousin’s (III.2) daughter, born in 2020, has early-onset AD and was recently found to have severe neutropenia.

In the index patient, laboratory findings were comprised of eosinophilia (up to 5900/µL [0-460]), elevated serum IgE up to max 1970 kU/L [<20] and decreased levels of IgG and IgM (Table 1).

A recent immunologic workup of the index patient revealed reduced NK cells, but normal T- and B-cell counts. Elevated levels of activated CD3+ T-cells, predominantly HLA-DR+CD8+ T-cells were observed, whereas the proportion of naïve T cells was normal. The proportion of naïve B-cells was elevated, but class-switched memory cells were significantly reduced (Table 1). Hypogammaglobulinemia and poor B-cell maturation were detected in the index patient only.

Other family members (II.1 and 2, III.1 and 2) also show increased IgE levels, activated HLA-DR+CD3+ T cells, effector memory CD45RA re-expressing CD8+ T-cells (TEMRA) and an elevated CD4/CD8 ratio, but markedly reduced T-cell function: decreased proliferation upon stimulation with IL-2, in a mixed lymphocyte culture and to specific antigens. Intracellular cytokine measurements indicated elevated IL-4 production in 2/3 patients tested, suggesting skewed Th2 differentiation (Table 1).

Whole-exome sequencing (WES) revealed a heterozygous variant in CARD11 (NM_032415.7: c.223C>T; p.Arg75Trp). This variant is absent from the gnomAD database, and is predicted to be damaging based on several in-silico algorithms (Polyphen2, SIFT MutationTaster), including a high CADD score of 29 (v1.6). The variant was also confirmed by Sanger sequencing in the mother, aunt and cousins of the index patient (Figure 1D). No other suspicious genetic variants were found by WES. The variant was submitted to the ClinVar database (SUB11100602).

Arg75 is a highly conserved amino acid residue located in the CARD domain, a region that is depleted of missense variation (Figure 1C and Supplementary Figure 2). Indeed, a recent multiplexed assay of variant function identified Arg75 as one of many hotspots for LOF variants in the CARD domain of CARD11 (13), and a separate amino acid exchange at this codon (p.Arg75Gln) was previously confirmed as DN in a patient with neutropenia (11). To formally test the impact of this novel variant on CARD11 signaling, we transfected JPM50.6 cells with expression plasmids encoding WT and/or R75W CARD11 and measured NF-κB driven GFP expression following T-cell activation (3). Relative to WT, no appreciable NF-κB induction was noted with R75W CARD11 (Figure 4A). Moreover, R75W demonstrated mild DN activity in suppressing NF-κB activation when co-expressed with WT CARD11 (Figure 4B), with comparable expression of both proteins noted. These results confirmed R75W as a causative DN CARD11 variant in this family.

DISCUSSION

This multi-generation family has clinical and immunological features in common with formerly described patients suffering
TABLE 1 | Summary of clinical and immunological characteristics.

|                          | Patient characteristics | Bacterial infections | Viral infections | Bronchiectases | Asthma bronchiale | Atopic dermatitis | Food and pollen allergies | Malignancies | Treatment |
|--------------------------|-------------------------|----------------------|------------------|----------------|------------------|-------------------|--------------------------|--------------|-----------|
|                          | Index patient III.4     | Patient’s mother II.2| Patient’s aunt II.1| Patient’s cousin f. III.2 | Patient’s cousin m. III.1 | Patient’s grandfather I.1 |
| **Age of onset**         | 2 y                     | early childhood      | 14 y             | early childhood | early childhood | early childhood     |
| **Bacterial infections** | Otitis media, sinusitis| n                    | n                | n              | n                | NA                |
| **Viral infections**     | HSV rectally            | p                    | n                | n              | n                | NA                |
| **Bronchiectases**       | y                       | n                    | y                | y              | y                | NA                |
| **Asthma bronchiale**    | y                       | y                    | y                | y              | y                | NA                |
| **Atopic dermatitis**    | y                       | y                    | y                | y              | y                | NA                |
| **Food and pollen allergies** | n                  | y                    | y                | y              | y                | NA                |
| **Malignancies**         | Anal squamous cell carcinoma | n                    | n                | n              | Mycosis fungosis, suspected T-cell lymphoma with lung metastases | NA |

|                          | Patient characteristics | Treatment |
|--------------------------|-------------------------|-----------|
|                          | Blood count             | Alive     |
| **Total leukocytes**     | 6600 [4400-8100]        | y         |
| (cells/µL)               |                        | (deceased at age 50) |
| **Eosinophils**          | 5900 [0-650]            | y         |
| (cells/µL)               | 130 [0-490]             | (deceased at age 56) |
| **Neutrophils**          | 494 [165-798]           | y         |
| (cells/µL)               | 342 [180-755]           | n         |
| **Lymphocytes**          | 1980 [1400-3300]        | y         |
| (cells/µL)               | 750 [1400-2800]         | y         |
| **Lymphocyte subsets**   |                         | n         |
| **CD4+ (cells/µL)**      | 1663 [1000-2200]        | y         |
|                         | 495 [1000-1980]         | (deceased at age 56) |
| **CD4+ (cells/µL)**      | 1228 [530-1300]         | y         |
|                         | 360 [630-1120]          | (deceased at age 56) |
| **CD8+ (cells/µL)**      | 396 [330-920]           | y         |
|                         | 128 [240-700]           | (deceased at age 56) |
| **CD4/CD8 ratio**       | 3.1 [1-2.5]             | y         |
|                         | 2.8 [1.3-3.5]           | (deceased at age 56) |
| **NK cells**             | 75 [90-700]             | y         |
| (CD3-CD16+CD56+ cells/µL) | 51 [80-630]         | (deceased at age 56) |
| **B cells**              | 238 [110-570]           | y         |
| (CD3+CD19+ cells/µL)     | 188 [90-320]            | (deceased at age 56) |
| **transitional B cells** | 4.9 [0.9-5.7]           | y         |
| (IgM+ +/CD38++ of %CD19+) | 3.6 [0.9-5.7]         | (deceased at age 56) |
| **naive B cells**        | 92 [52-87]              | y         |
| (IgD+CD27+ of %CD19+)    | 80 [52-87]              | (deceased at age 56) |

Continued...
### TABLE 1  | Continued

|                          | Index patient III.4 | Patient’s mother | Patient’s aunt | Patient’s cousin f. III.2 | Patient’s cousin m. III.1 | Patient’s grandfather |
|--------------------------|----------------------|------------------|----------------|--------------------------|---------------------------|------------------------|
| class-switched (IgM-/CD20+/CD27+ of %CD19+) |                      |                  |                |                          |                           |                        |
| Th1 (% of IFNγ expression in CD3+) | 2,7 [3,8-23]         | 4,7 [3,8-23]     | 7,7 [3,8-23]   | 5 [3,8-23]               |                           | 14,1 [3-46]            |
| Th2 (% of IL-4 expression in CD3+) | 0,75 [0,12-0,78]     | NA               | 1,91 [0,12-0,78] | [3,8-23] 3,21 [1,12-0,78]|                           | NA                     |
| Th17 (% of IL-17 expression in CD3+) | 3,12 [1,41-4,73]     | NA               | 1,94 [1,41-4,73]| 4,72 [1,41-4,73]          |                           | NA                     |
| T-cell proliferation     |                      |                  |                |                          |                           |                        |
| IL-2                     | ↓                    | ↓                | ↓              | normal                   |                           | NA                     |
| PHA                      | normal               | normal           | normal         | normal                   | normal                   | NA                     |
| CD3/CD28 beads           |                      |                  |                |                          |                           |                        |
| MLC pool                 | ↓                    | ↓                | ↓              | ↓                        | NA                       | NA                     |
| Tetanus antigen          | ↓                    | ↓                | ↓              | ↓                        | ↓                        | NA                     |
| CMV antigen              | ↓                    | ↓                | ↓              | ↓                        | NA                       | NA                     |
| Adenovirus antigen       | normal               | normal           | ↓              | ↓                        | ↓                        | NA                     |
| Candidin antigen         | ↓                    | ↓                | ↓              | ↓                        | NA                       | NA                     |
| Immunoglobulins          |                      |                  |                |                          |                           |                        |
| IgG (g/L)                | 7,1¹ [7-16]          | 6,9 [7-16]       | 8,43 [7-16]    | 8,64 [7-16]              |                           | 11,89 [7-16]           |
| IgM (g/L)                | <0,16 [0,4-2,3]      | <0,06 [0,4-2,3]  | 0,7 [0,4-2,3]  | 0,37 [0,4-2,3]           | 0,72 [0,4-2,3]           | NA                     |
| IgE (kU/L)               | 1970 [<20]           | <2 [<20]         | 1973 [<20]     | 4798 [<20]               |                           | 575 [<20]              |
| IgA (g/L)                | 2,07 [0,7-4]         | 1,48 [0,7-4]     | 2,22 [0,7-4]   | 4,36 [0,7-4]             |                           | 6,39 [0,7-4]           |

¹Under immunoglobulin substitution; ²Sample taken after radiotherapy; normal >30% of healthy control; MLC, mixed lymphocyte culture; NA, not available; y, yes; n, no; ↓ - reduced; bold values – abnormal results.
from CADINS disease (1, 8). As described for other CARD11 DN variants, there is a high penetrance but variable intrafamilial expressivity caused by the (p.Arg75Trp) variant. As an infant, the index patient had atopic dermatitis-like rash, but was more severely affected by infections and early-onset asthma. Other family members suffered from moderate to severe atopic symptoms that raised no suspicion for an IEI. The agranulocytosis of the index patient was detected at three months of age, after one year the neutrophil count had normalized. Dorjbal et al. observed neutropenia in 4 patients with functional DN variants not associated with bone marrow abnormalities suggesting an autoimmune origin (1).

Although T-cell subpopulations were normal in numbers and distribution, the proliferation of the index patient’s T cells was impaired upon stimulation with mitogens such as IL-2 and in mixed lymphocyte culture, but also after stimulation with specific antigens such as tetanus, CMV and candidin. This was also observed in all other affected family members and is consistent with impaired CARD11-dependent T-cell activation after TCR (CD3) and CD28 ligation. Indeed, impaired T-cell function predisposes to viral infections, and poor CD28 stimulation seems to have a special significance in the predisposition for HPV infection (14, 15). Apart from CARD11 deficiency, signaling via CD28 is also disturbed in other IEI such as CD28, MAGT1 and CARMIL2 deficiencies, and affected patients have been documented with HPV-driven warts (1, 16, 17). In a cohort of 44 individuals with CADINS disease, cutaneous infections were the second most common manifestation (68%) after AD (73%). Viral skin infections included common HPV-driven warts in over 25% of patients with CARD11 DN variants (1). HPV is known to cause malignancies like cervical cancer and is linked to more than 90% of all anal carcinomas (18) e.g. in HIV positive individuals. These findings highlight a possible relationship between this germline CARD11 variant and the HPV18 positive anal carcinoma in the index patient’s mother. HPV16 has been identified as the predominant HPV type in anal squamous cell carcinoma, followed to a lesser extent by HPV18, HPV33, HPV31, HPV6, HPV58 and HPV35 (19). HPV vaccination which is 9-valent (HPV types 6, 11, 16, 18, 31, 33, 45, 52, and 58) may offer an option for primary prevention of HPV induced carcinomas in patients with CADINS in analogy to HIV patients. Likewise, routine screening including anoscopy and anal cytology may be warranted in CADINS patients (16, 18).

Somatic CARD11 variants have been associated with several lymphoid malignancies (20). Somatic GOF CARD11 mutations were initially described in ~10% of diffuse large B-cell lymphomas associated with constitutive NF-kB activity (21). Moreover, hypermorphic/GOF variants in CARD11 and other non-canonical NF-kB genes were also reported in patients with MF, Sézary Syndrome, and HTLV-1-driven adult T-cell lymphoma (20, 22, 23). More recently, somatic variants in CARD11 were found to be associated with solid tumors such as human sinonasal tumors (24). Watt et al. showed that in cutaneous squamous cell carcinomas, CARD11 protein expression is increased, and novel CARD11 variants outside the coiled-coil domain led to constitutively activated NF-kB signaling (25). By contrast, germline CARD11 LOF/DN variants were previously discovered in 2 patients with peripheral T-cell lymphoma (p.Glu57Asp) and MF (p.Arg187Trp) (11). Our findings further underscore a perplexing connection between attenuated CBM signaling and malignancy, perhaps linked to infectious triggers (e.g. HSV-1) and/or impaired T-cell surveillance in the skin. More work is required to decipher how defects in distinct CARD11-dependent signaling pathways may explain this surprising phenomenon.

Dupilumab, a humanized monoclonal antibody against IL-4Rα, is increasingly being used in treatment-refractory AD, including in children from the age of 6 years. The imbalance towards Th2 cytokine production in CARD11 DN patients suggests that this biological may be useful in controlling AD in IEI with atopy. Recently, it has been successfully administered in
the treatment of AD in monogenic forms of HIES, such as STAT3 LOF (3) and also in one individual with a DN CARD11 variant (26). Moreover, a recent meeting report described three patients with CADINS disease and severe AD ameliorated by treatment with dupilumab oromalizumab (27).

In summary, we describe a multi-generation family with CADINS disease caused by a novel CARD11 DN variant. The clinical phenotype is not only characterized by severe atopy (AD, asthma, food allergies) and recurrent infections in the context of both hypogammaglobulinemia and reduced T-cell responses to specific antigens, but also through the occurrence of 2 different malignancies: an HPV-driven anal squamous cell carcinoma and skin infiltrating T-cell lymphoma in one family member each. Dupilumab is likely helpful in alleviating AD in some CADINS patients.

METHODS

Patients were enrolled on protocols approved by ethics review committees and thereupon provided written informed consent.

Immunophenotyping

Patient peripheral blood mononuclear cells (PBMC) were isolated by density-gradient centrifugation using Biocell separating solution (density 1.077 g/ml, Biocell) and isolated by density-gradient centrifugation using Biocell. Patient peripheral blood mononuclear cells (PBMC) were used, applying BD FACS Lyse Wash Assistant, for B-cell immunophenotyping in the following manner: for T-cell panel - 100µl blood (EDTA) were used, applying BD FACS Lyse Wash Assistant, for B-cell panel - BD Vacutainer CPT 4ml was used, preparation of peripheral blood mononuclear cells (PBMCs) was performed according to manufacturer’s protocol. Antibodies used: CD45, CD14, CD19, CD20, CD27, CD16/56, CD3, CD4, CD8, TCRab, CD3/28, HLA-DR, CD45RA, CD45RO, CCR7, CD31, CD21, CD14, CD19, CD20, CD27, CD16/56, CD3, CD4, CD8, TCRab, CD3/28. Samples were acquired and analysed with BD FACSlyric Flow Cytometer using the BD FACSuite V1.4.0 Analysis-Software.

Intracellular Cytokine Staining

Intracellular cytokine measurement was performed according to Körholz et al., 2021 (7). Briefly PBMCs (1x10⁶/ml) were left unstimulated or stimulated with 10 ng/ml PMA (Phorbol-12-Myristat1-13-Aacet) and 1µg/ml Calcium-Ionophore (both Merck KGaA, Darmstadt, Germany) under addition Brefeldin A (1µl/ml, BD-Biosciences San José, CA) for 12h overnight. Cells then were harvested and washed twice with PBS/1%FCS. For surface staining, cells were incubated with anti-CD45RO-PE (5µl/test) (Biolegend, San Diego, CA), anti-CD3-APC (2µl/test), anti-CD4-APC-AlexaFluor750 (5µl/test) and anti-CD45-Krome Orange (5µl/test) (Beckman Coulter, Krefeld, Germany) for 30 min at 4°C. After being washed twice with PBS/1% FCS, cells were fixed and permeabilized with 100µl Cytofix/Cytoperm™ (BD Biosciences, San José, CA) for 20 min at 4°C and then washed twice according to the manufacturers instructions. For intracellular staining, cells were incubated with anti-IFNy-FITC (5µl/test), anti-IL4-PE Cy7 (5µl/test) (both Biolegend) and anti-IL-17A eFlour 450 (5µl/test) (eBioscience/Thermo Scientific) and anti-CD4 APC-AlexaFluor750 (2µl/test) (Beckman-Coulter, Krefeld,Germany) for 45min at 4°C. Cells were washed twice with Perm/Wash Buffer (BD Biosciences) and diluted in 500µl PBS/1% FCS. Analysis was performed on a BD LSR II.

T-Cell Proliferation

T-cell functions were assessed using a standard 3-H-Thymidin proliferation assay: 10⁵ cells/well were stimulated in triplicates in a 96 well flat bottom plate with IL2 (100IE/well, Pepro Tech, New Jersey, US), PHA (1,5%, Invitrogen, Carlsbad, CA, USA), CD3/28 beads (12,5µl/ml, Thermo Scientific/Gibco Waltham, MA, USA), Tetanus antigen (50µl/ml, Statens Serum Institut, Copenhagen, Denmark), CMV antigen, Adenovirus antigen (both: Institute for Virology, Ulm University). As a positive control, buffy coat PBMC of healthy donors were stimulated. After 3 days of incubation for unspecific stimulation (IL2, PHA, CD3/28) and 5 days for antigen-specific stimulation (Tetanus, viral antigens) at 37°C/5% CO₂, ³H-thymidin (0,001mCi/ml, Perkin Elmer Germany) was added for 18h. Cells were harvested and incorporated ³H-thymidin was measured in a beta counter (Microplatecounter TopCount NCT, Packard/Perkin Elmer). Stimulation indices were calculated as cpm mean (of triplicates) after stimulation/cpm mean of negative controls (culture-medium alone).

Molecular Genetics Analysis

Genomic DNA from the index patient was isolated from peripheral blood and coding genes enriched using the SureSelect Human All Exon Kit V6 (Agilent technologies) for subsequent WES on the Illumina system. Reads were aligned to the human genome build GRCh37/hg19. Sequence reads were called and analyzed according to an in-house standard operating procedure using the VarFish platform (28). Variants were filtered by minor allele frequency, mode of inheritance, functional prediction and constraint metrics, such as LOEUF score and pLI score for LoF-variants and Z-Score for Missense variants (29). Confirmation of the CARD11 variant and segregation analysis was done by Sanger sequencing using the following primers for amplification and sequencing: Ex4.F:5’-GACCTTGGTCTCCATCGATATGT-3’, Ex4.1R:5’-ACACGCCCTCCTCCTTAGAG-3’.

Cell Transfections

pUNO-CARD11-FLAG plasmids encoding the R75W CARD11 protein were produced by site-directed mutagenesis as previously described (11). JPM50.6 T cells (3 million) were transfected with 3 µg empty pUNO vector (EV), WT or R75W CARD11 plasmids, with or without WT CARD11-V5 using an ECM 630 electroporator (Harvard BTX, Holliston, MA, USA) at 260V, 950 µF (11). At 24 hrs post-transfection, cells were stimulated for 24 hrs with 1 ng/ml of anti-CD3 (clone HIT3a) + anti-CD28 (clone CD28.2) antibodies (BD Biosciences), 20 ng/ml phorbol myristate acetate (PMA) + 1 mM ionomycin (Millipore Sigma, St. Louis, MO, USA) or left unstimulated. NF-kB-dependent
GFP reporter expression was measured using an Accuri C6 flow cytometer (Becton Dickinson). CARD11 protein expression was assessed in transfectant lysates by immunoblotting as previously described (11).

**Histology and Molecular Pathology**

The index patient’s mother had a recurrent moderately well differentiated squamous cell carcinoma of the anal canal which was operated and routinely processed. The postoperative tumor classification was: rPT4, rNx, rL0, rV0, rPn0, Rx, G2. **Figure 3** shows a different magnification of the H&E sections and a strong and diffusely positive immunohistochemical reaction for p16 (Cytomed mouse monoclonal antibody clone JC2; 1:100, OPTIview detection system on a VENTANA platform) which was used as a surrogate marker for HPV-infection.

DNA was extracted from the tumor sample for HPV-PCR and subsequent typing by means of the HPV 3.5 LCD-array Kit (Chipron). Typing revealed infection with the carcinogenic HPV-subtype HPV18.

**CONCLUSION**

Patients with CARD11-associated atopy and dominant interference of NF-kB signaling (CADINS) disease, were first described in 2017. They usually present with varying degrees of atopy and cutaneous viral and respiratory infections with or without hypogammaglobulinemia. Our study expands the clinical spectrum of CADINS patients by reporting a multi-generation family with Hyper-IgE, disturbed T-cell function in response to IL-2, MLC and specific antigens, hypogammaglobulinemia and malignancies. Poor CD28 signaling may explain predisposition to HPV infection. An HPV triggered squamous cell carcinoma has, however, not been reported in CARD11 deficiency before. This finding underscores a perplexing connection between attenuated CBM signaling and malignancy, perhaps linked to infectious triggers and/or impaired T-cell surveillance. Further investigation is warranted to elucidate how defects in distinct CARD11-dependent signaling pathways may explain this surprising phenomenon. Dupilumab is likely helpful in alleviating severe AD in some CADINS patients.

**DATA AVAILABILITY STATEMENT**

The datasets for this article are not publicly available due to concerns regarding participant/patient anonymity. Requests to access the datasets should be directed to the corresponding author.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethikkommission an der TU Dresden. The patients/participants provided their written informed consent to participate in this study. Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

**AUTHOR CONTRIBUTIONS**

LP and JK, conceptualization, data collection, writing, and editing. FB, formal analysis (interpretation of exome data), software, and visualization. MS, methodology and HPV sequencing. DP, technical assistance and methodology. EK, patient care, methodology, and photodocumentation. CK, ML, and RB, patient care and editing. EJ, data curation and investigation. JR, methodology, writing, editing, and treatment of the index patient. BD, DY, and AS, investigation, functional testing of the variant, and editing. DA, methodology, data interpretation, and editing. VG and ML-K, Sanger sequencing, editing, and funding acquisition. CS, conceptualization, data curation, writing-editing, supervision, and funding acquisition. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.878989/full#supplementary-material

**Supplementary Figure 1** | Structural modeling of the CARD11 protein by AlphaFold (https://alphafold.ebi.ac.uk/entry/Q9BXL7). Positively charged amino acid Arg75 forms hydrogen bonds with negatively charged amino acid Glu59, likely important for protein stability of the CARD domain.

**Supplementary Figure 2** | Screenshot from DECIPHER - https://www.deciphergenomics.org/gene/CARD11/overview/protein-genomic-info - showing domain structure, regional missense constraint scores and ClinVar annotated pathogenic variants for the CARD11 gene. Notably, amino acid residues 1-140 (including the CARD domain) are depleted of missense variation (calculated “MPC” of 0.1 indicated by black asterisk – data retrieved from ExAC). Pathogenic missense variants cluster within the CARD domain. Black triangle points towards the location of the variant described in this study (p.Arg75Trp).
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