Constitutional SAMD9L mutations cause familial myelodysplastic syndrome and transient monosomy 7


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Constitutional SAMD9L mutations cause familial myelodysplastic syndrome and transient monosomy 7

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ABSTRACT

Familial myelodysplastic syndromes arise from haploinsufficiency of genes involved in hematopoiesis and are primarily associated with early-onset disease. Here we describe a familial syndrome in 7 patients from 4 unrelated pedigrees presenting with myelodisplastic syndrome and loss of chromosome 7/7q. Median age at diagnosis was 2.1 (1-42) years, all patients presented with thrombocytopenia with or without additional cytopenias and a hypocellular marrow without increase of blasts. Genomic studies identified constitutional mutations (p.H880Q, p.R986H, p.R986C and p.V1512M) in SAMD9L gene on 7q21, with decreased allele frequency in hematopoiesis. The nonrandom loss of mutated SAMD9L alleles was attained via monosomy 7, deletion 7q, UPD7q, or acquired truncating SAMD9L variants p.R1188X and p.S1317RfsX21. Incomplete penetrance was noted in 30% (3/10) of mutation carriers. Long-term observation revealed divergent outcomes with either leukemic progression and/or accumulation of driver mutations (n=2), persistent monosomy 7 (n=4), and transient monosomy 7 followed by spontaneous recovery with SAMD9L-wildtype UPD7q (n=2). Dysmorphic features or neurological symptoms were absent in our patients, pointing to the notion that myelodisplasia with monosomy 7 can be a sole manifestation of SAMD9L disease. Collectively, our results define a new subtype of familial myelodysplastic syndrome and provide an explanation for the phenomenon of transient monosomy 7. www.clinicaltrials.gov; #NCT00047268
INTRODUCTION

Germline predisposition has been recognized as an underlying cause for the development of myelodysplastic syndromes (MDS) in children. Recently, it has also been gaining importance in the etiology of adult myeloid neoplasia (MN), particularly in cases with a positive family history. In addition to inherited bone marrow failure syndromes (IBMFS), a number of genes are known to be associated with heritable forms of MDS and acute myeloid leukemia (AML)(1, 2), i.e. GATA2(3), CEBPA(4), RUNX1(5), ANKRD26(6), ETV6(7) and DDX41(8). Germline mutations in DDX41 can result in adult onset MN, while aberrations in RUNX1 and GATA2 are associated with MN in younger individuals. We recently reported that GATA2 deficiency is the most common genetic cause of primary childhood MDS, accounting for 15% of all advanced MDS cases, and 37% of MDS with monosomy 7 (MDS/-7)(9). However, in the majority of pediatric MDS, and also in a considerable number of familial MN cases, the presumed germline cause has not yet been discovered(10) (11).

Monosomy 7 is the most frequent cytogenetic lesion in children with MDS and, unlike in adults, it often occurs as the sole cytogenetic abnormality(12). Due to the rapid and progressive disease course it is considered an urgent indication for hematopoietic stem cell transplantation (HSCT)(13). However, transient monosomy 7 has been occasionally documented in childhood MDS(14-16). Considering that complete (-7) and partial (del(7q)) deletion of chromosome 7 are common aberrations in MDS, extensive efforts have been undertaken to discern causative tumor suppressor genes located on chromosome 7. Asou and colleagues identified SAMD9 (Sterile Alpha Motif Domain-containing 9), its parologue SAMD9L (SAMD9-like), and Miki/HEPACAM2 as commonly deleted genes within a 7q21 cluster in patients with MN(17).
Notably Samd9l-haploinsufficient mice were shown to develop myeloid malignancies characterized by different cytopenias and mimicking human disease with monosomy 7(18).

In line with these findings, germline heterozygous gain-of-function (GoF) SAMD9L mutations p.H880Q, p.I891T, p.R986C, and p.C1196S were recently discovered in 4 pedigrees with variable degrees of neurological (ataxia, balance impairment, nystagmus, hyperreflexia, dysmetria, dysarthria) and hematological (single to tri-lineage cytopenias, MDS/-7) symptoms. For most carriers, the clinical presentation was compatible with the diagnosis of ataxia-pancytopenia (AT) syndrome(19, 20). Similarly, in two recent studies, we and others reported de novo GoF mutations in SAMD9 in a total of 18 patients with MIRAGE syndrome (myelodysplasia, infection, restriction of growth, adrenal hypoplasia, genital phenotypes, and enteropathy), of whom 4 notably also developed MDS/-7(21, 22), however not all patients develop the full MIRAGE disease spectrum, as documented in one family with SAMD9-related MDS(23). SAMD9 and SAMD9L genes share 62% sequence identity and apart from their putative role as myeloid tumor suppressors, the general function and their specific effect pertaining to hematopoiesis are not well-understood(18).

In this study we aimed to identify the genetic cause in pedigrees with nonsyndromic familial MDS. We discovered constitutional SAMD9L mutations associated with nonrandom patterns of clonal escape leading to loss of the mutant allele. We further demonstrate in two cases that SAMD9L–related disease can be associated with transient -7, occurring as a one-time clonal event followed by somatic correction of hematopoiesis achieved by UPD7q with double wild type SAMD9L.
METHODS

Patients

Diagnosis of MDS was established according to the WHO criteria(24, 25). Patients 1 (P1), 2 (P2), 5 (P5), and 7 (P7) were enrolled in the prospective study 98 of the European Working Group of MDS in Childhood (EWOG-MDS) (www.clinicaltrials.gov; #NCT00047268). Patient 6 (P6) was the father of P5 (family III). Family II (P3 and P4) was referred for evaluation of familial MDS from the Phoenix Children’s Hospital, USA. The study had been approved by the institutional ethics committee (University of Freiburg, CPMP/ICH/135/95 and 430/16). Written informed consent had been obtained from patients and parents.

Genomic studies and bioinformatics

Exploratory whole exome sequencing (WES) was performed in bone marrow granulocytes (BM-GR) in P1 and P2, as outlined in the supplement. Targeted deep sequencing (DS) for $\text{SAMD9/SAMD9L}$ and MDS/IBMFS-related genes was performed in other patients, except for P3 and P6 due to unavailability of material. All relevant variants were validated using Sanger sequencing. For germline confirmation, DNA was extracted from skin fibroblast and/or hair follicles, and targets were amplified and sequenced as previously described(9). The degree of deleteriousness was calculated using the combined annotation-dependent depletion scoring system (CADD-score)(26). The variants with CADD-scores higher than 20 were further evaluated towards their role in hematological disease or cancer, thereby focusing on the top 1% most deleterious variants in the human genome. In addition, the pathogenicity calculations were achieved using standard prediction tools. The evolutionary conservation across species and the
physicochemical difference between amino acids was estimated by PhyloP, PhastCons and Grantham score respectively(27). Mutant clonal size was inferred from allelic frequencies and the total number of NGS reads normalized to the ploidy level. Further details are provided in supplementary methods.

**Evaluation of variant allelic configuration**

Genomic DNA of P1 collected at the time point of progress to chronic myelomonocytic leukemia (CMML) was amplified to obtain a 1333 bp region encompassing both \( \text{SAMD9L} \) mutations p.V1512M (germline) and p.R1188X (acquired). PCR products were TA-cloned and sequenced as previously reported(28). Sequences of 170 colonies were evaluated for the presence of \( \text{SAMD9L} \) mutations.

**Cellular and functional studies**

Metaphase karyotyping and interphase fluorescence in situ hybridization (FISH) were performed using bone marrow (BM) specimens according to standard procedures(12). Human colony forming cell (CFC) assays were performed in P1 (at CMML disease stage) and in P7 (at diagnosis) as previously described(29). Further, to evaluate the effect of the patient-derived \( \text{SAMD9L} \) p.V1512M and p.R986C mutations on cellular proliferation, 293FT cells were dye-labeled and consequently transfected with wild type or mutant teal fluorescent protein (TFP)-\( \text{SAMD9L} \) as previously described(20). The transfected cells were tracked by flow cytometry for TFP-\( \text{SAMD9L} \) expression and dye dilution as an indicator of cell division. \( \text{SAMD9L} \) variant p.T233N recently reported as “disease-protective” (20) was used as control.
RESULTS

Clinical phenotype of patients

Patients P1-P6 (4 males and 2 females) belong to 3 unrelated families of German descent and were diagnosed with bona fide familial MDS after the exclusion of known IBMFS achieved by targeted sequencing and functional tests (Figure 1, Table 1). Index patient P7 is the only child of a non-consanguineous Swedish family. Detailed clinical description of all patients is outlined in the supplement. All affected individuals had normal measurements without dysmorphic stigmata at birth and at last follow-up (Table 1). Psychomotor development and neurocognitive function were normal, in particular no ataxia or movement disorder were diagnosed in the 10 mutation carriers (7 patients and 3 silent carriers with SAMD9L mutations). Previous family histories were unremarkable for cytopenias, neurological disease, malignancies, or stillbirths, with the exception of father of P7 who at the last follow-up presented with unclear ataxia. Prior non-invasive recurrent respiratory tract infections were noted in 3/7 patients (P1, P2, and P4) and endogenous eczema in 2 (P1 and P3). Moreover, P1 developed transient pancytopenia during infancy and suffered from self-limiting seizures during infancy with no structural brain abnormalities or neurological deficits identified. Peripheral blood findings at diagnosis included isolated thrombocytopenia in one (P2), thrombocytopenia with neutropenia in two (P1, P3) and pancytopenia in 4 patients (P4-7). MCV was elevated in 5/7 patients at diagnosis, and HbF in 2 out of 3 tested patients. MDS manifested at the median age of 2.1 (1.0-42) years as hypocellular RCC or in the adult (P6) refractory cytopenia with multilineage dysplasia (RCMD, Table 1). Severe dysplasia with vacuolization was observed in 2 patients (P1-2). A common cytogenetic feature in all patients was the complete or partial loss of chromosome 7 (Table 1).
The clinical course was remarkable for several patients. P1 developed 3 years after the initial diagnosis of RCC severe infections and hepatosplenomegaly, and for the first time required platelet transfusions, while his blood smear depicted 21% blasts compatible with the diagnosis of CMML (with no in vitro GM-CSF hypersensitivity), (Figure 2). Following HSCT, he developed late-onset acute GVHD and died from cerebral hemorrhage. At the same time, his younger sister (P2) was diagnosed with RCC/-7, however the parents decided against follow-up in hematology clinic. Unexpectedly, her complete blood count (CBC) normalized 3.7 years later and remained stable until last follow-up 20 years after the initial diagnosis. She achieved a spontaneous remission as supported by normalization of BM morphology/cellularity and cytogenetics (Figure 2, Table 1). A similar clinical picture was seen in P7 who was diagnosed with RCC with a -7 clone at the age of 2.1 years, and experienced rapid cytogenetic remission with normal marrow and blood counts until the last follow-up, 16 years after diagnosis (Table 1). In P4, shortly after initial manifestation, hematology normalized, however FISH revealed chromosome 7 loss in BM, which gradually increased and after 3.5 months culminated in the emergence of two independent clones with -7 and del(7q) (Table 1).

Due to high-risk cytogenetics and disease progression, 5/7 patients (P1, P3-6) underwent HSCT after myeloablative conditioning. At last follow-up, 6 out of 7 patients were alive, 4 after HSCT, and 2 without therapy.

**Constitutional and acquired **SAMD9L** mutations**

Exploratory WES performed in family I identified two shared candidate variants in P1 and P2 evaluated as highly conserved and deleterious by in silico prediction: **SAMD9L** (p.V1512M) and
PTEN (p.Y188C), (Table 2, Figure S1). Sequencing of DNA from hair follicles confirmed the constitutional nature of both novel mutations. SAMD9L p.V1512M variant was inherited from the mother (Figure 1A) whereas PTEN p.Y188C was of paternal origin; both parents were asymptomatic and had normal CBC at the time of testing. Finally, truncating acquired SAMD9L mutation p.R1188X (VAF 5.9%) was identified in P1 in hematopoiesis (Table 2).

In pedigree II, targeted NGS in P4 revealed SAMD9L p.R986H as the most plausible candidate constitutional mutation predicted to be highly conserved and deleterious (Table 2, Figure 1A-B). This mutation was found in 4 individuals in ExAC (out of 120976 alleles). Additional missense variants in JAK3 p.A877V (ExAC: 11 individuals, 121372 alleles), and FANCM p.L57F (ExAC: 195 individuals, 121190 alleles) had lower and moderate pathogenicity scores, respectively (Table 2). Chromosomal breakage studies on P4 were negative thus arguing against a pathogenic role of the heterozygous FANCM variant. Germline analysis revealed SAMD9L and JAK3 variants in P3, P4, and their father, while the FANCM variant was transmitted from the mother only to P4. Both parents were asymptomatic.

In pedigree III, the SAMD9L p.R986C mutation was identified in P5 and the affected father, P6 (Figure 1A). This mutation has been reported in a family with ataxia-pancytopenia phenotype, with 1 of 3 carriers developing MDS/-7 at the age of 18 months(20). The HLA-identical brother of P5 was thoroughly evaluated as a potential BM donor. He was clinically healthy and had normal CBC, but he did not qualify as a donor because of hypocellular BM with mild dysplastic features. He was also carrier of the p.R986C mutation.
In P7 of pedigree IV, targeted NGS identified two \textit{SAMD9L} mutations (Table 2): missense p.H880Q with VAF of 27% out of 8139 reads (likely constitutional; this mutation was reported in multiple individuals within a family with ataxia pancytopenia but no MDS phenotype) and nonsense p.S1317RfsX21 likely acquired in a subclone as inferred from the much lower VAF of 10% (5934 reads). In summary, inherited \textit{SAMD9L} mutations p.V1512M, p.R986H, p.R986C were identified in 3 families (each with 2 individuals diagnosed with MDS/-7 and 1 healthy carrier, Figure 1A-B), and p.H880Q in P7 who presented with transient monosomy 7.

\textbf{Acquired mutations in known oncogenes}

All patients with exception of P3 and P6 were evaluated for the presence of somatic mutations in leukemia-associated genes using WES or targeted NGS. In P1 previously reported leukemia driver mutations \textit{SETBP1} p.D868N, \textit{EZH2} p.V582M, \textit{KRAS} p.Q61P were identified as somatic (Table 2, Figure S2). Similarly, in P5 a somatic \textit{RUNX1} mutation c.413_427+5dup20bp was detected (Table 2). This is a novel splice-donor site mutation with no occurrence in databases. No additional mutations in leukemia driver genes were observed in other affected cases.

\textbf{Clonal escape mechanisms from \textit{SAMD9L} mutations are not random}

In comparison with other constitutional variants, \textit{SAMD9L} missense mutations showed significantly lower median allelic frequencies across all patients (53% vs. 20%, p<0.05, Table 2). This finding was corroborated by the consistent partial or complete loss of chromosome 7 (Figure 3, Table 1). In P1, -7 progressively expanded from 60% at time of RCC diagnosis to 95% at CMML progression. In addition, P1 and P7 harbored subclones with acquired stop-gain
SAMD9L mutations located upstream of the constitutional missense substitutions (Figure 3). TA cloning confirmed cis orientation of both mutations on the same chromosome in P1 (Figure 1C). Western blot of 293T cells transiently transfected with double-mutated (p.V1512M/p.R1188X) SAMD9L plasmid revealed stable expression of truncated protein (not shown). In P2 and P7, the initial −7 disappeared and was replaced by a large somatic clone with uniparental disomy (UPD) 7q that duplicated the paternal wildtype SAMD9L allele (Figure 3-4).

**Natural history of SAMD9L-related MDS reveals divergent clinical outcomes**

We next studied in detail the disease course in patients who were left untreated for longer periods of time. In P4, the evolution from normal karyotype to independent clones with −7 and del(7q) was rapid and occurred within a few months (Table 1). Cytogenetic progress was associated with partial recovery of CBC and normalization of BM cellularity, pointing to the possibility of HSC niche repopulation by −7/del(7q) retaining only the wildtype SAMD9L allele. P1 developed CMML 3.6 years after initial diagnosis of MDS/−7. This patient carried somatic driver mutations representing major subclones co-existing within −7 background (Table 2). Additional somatic SAMD9L mutation p.R1188X (co-occuring in cis- with p.V1512M) was found in a minor clonal fraction of ~6%.

In contrast, P2 and P7 showed an unexpected clinical course with spontaneous hematologic recovery, disappearing monosomy 7 and the presence of large double wildtype UPD7q clone in BM (Figure 4-5). Both patients remained healthy, had normal follow-up BM examinations with no signs of dysplasia (Table 1), and normal CBC until last FUP, 20 (P1) and 16 (P7) years after initial diagnosis.
**SAMD9L mutations inhibit cell proliferation**

Inhibitory effect on cell proliferation reported for **SAMD9L** mutants overexpressed in 293FT cells *in vitro* was termed as gain-of-function (GoF). In contrary, ectopic expression of p.T233N was shown to mitigate cell proliferation to a smaller extent in comparison to WT **SAMD9L**, and was categorized as a disease-protective or loss-of-function (LoF) variant(20). **SAMD9L** p.R986H was previously functionally studied and shown to be GoF (20), while p.H880Q was shown to induce LOH by -7/del(7q) or UPD7q in an EBV-transformed cell line *in vitro*(19). To determine the effect of the **SAMD9L** MT identified in families I and III, we transiently transfected 293FT cells with vectors containing disease-associated mutations p.V1512M and p.R986C respectively, along with the LoF variant p.T233N. Cell proliferation was assessed in dye dilution assays. Both **SAMD9L** p.V1512M and p.R986C decreased dye dilution in comparison to **SAMD9L** WT and p.T233N (Figure 6A-B), pointing to an amplified growth restrictive effect of the disease-associated variants (Figure 6B).
DISCUSSION

In this study, we describe a familial MDS syndrome caused by heterozygous missense mutations in *SAMD9L* gene located on chromosome 7q21. We present 7 individuals from 4 unrelated pedigrees who developed MDS/-7 from the age of 1 to 42 years without any neurological involvement. Compared to the reported *SAMD9L* mutation carriers (19, 20), hematology was the sole clinical manifestation in our cases. Other novel findings outlined here are the description of the somatic mutational landscape likely contributing to MDS progression, the observation of transient monosomy 7, and finally the occurrence of nonrandom revertant mosaicism leading to complete hematological recovery.

The *SAMD9L* mutations p.R986H, p.R986C, p.V1512M identified in this cohort affect evolutionary highly conserved amino acid residues, are assessed as pathogenic by *in silico* prediction. The mutation p.H880Q (P7) shows a weak conservation score, however, this mutation had already been reported as causative for the ataxia pancytopenia phenotype (19). We were not able to test *SAMD9L* genetics in P7, however the unclear ataxia this patient had been evaluated for at last visit points to a carrier status and indicates that there must be an overlap between sole hematological and ataxia phenotype in *SAMD9L* disease. Summarizing all *SAMD9L* mutations recently reported or identified in our cohort, a total of 6 germline mutations can be discerned (p.H880Q, p.I981T, p.R986H, p.R986C, p.C1196S, and p.V1512M). Of note, all these mutations cluster exclusively to the C-terminal half of the protein. Further, upon comparing reported mutations in the paralogue gene *SAMD9* (p.R982H/C)(22) with that of the present study in *SAMD9L* (p.R986H/C), we identified a potential mutational hotspot affecting...
highly conserved regions in both SAMD9L (p.984-989: GVRIIH) and SAMD9 (p.980-985: GVRIIH) proteins.

The reported constitutional variants in both SAMD9/SAMD9L were classified as GoF based on the observation of decreased cell proliferation in a 293T cell line (20-22). Similarly, we observed growth deficiency in 293T cells harboring SAMD9L p.V1512M and p.R986C. Based on these findings, one cautious speculation hints at a GoF effect that is cell toxic. This can be supported by the discovery of an acquired stopgain SAMD9L mutations in P1 and P7 that likely “eliminate” germline missense mutations. In the cases studied here, complete or partial deletion of chromosome 7 and also UPD7q was nonrandom and each time resulted in loss of the germline-mutated SAMD9L gene copy. Additional studies are essential to further define the effect of SAMD9L variants, might be challenging due to the growth inhibitory effect of the alterations. It also remains to be answered if SAMD9L missense mutations lead to an increased protein stability, alter the protein structure, enhance an unknown functional domain, or exert a completely neomorphic effect.

We describe 3 silent mutation carriers from separate families demonstrating no previous medical history. Despite normal CBC and MCV, the brother of P5 exhibited a hypocellular marrow with mild dysplasia, evidently attributed to the identical pathogenic SAMD9L mutation. This finding emphasizes the necessity of thorough hematological workup including marrow studies in potential sibling donors especially when they lack a genetic marker for familial disease. The intrafamiliar heterogeneity regarding the hematologic presentation remains
elusive; one can speculate that other yet unknown genetic or epigenetic mechanisms might act as modifiers.

Thus far only limited knowledge exists about the regulation and cellular functions of *SAMD9L*. It has been postulated that both *SAMD9L* and the adjacent paralogous *SAMD9* gene share functional redundancy, shaped by a long term, possibly virus-induced selective pressure(30). Both genes can be upregulated by type I and II interferons(20, 31-33) and by this might suppress inflammatory pathways and exert anti-viral properties(31, 34). Although there was no evidence for a defined immunodeficiency in our cohort, 3/7 patients experienced recurrent respiratory infections with or without cytopenias before they developed RCC/-7.

Further evidence supports that *SAMD9/SAMD9L* genes act as tumor suppressors as their inactivation is associated with increased cellular proliferation, i.e. in normophosphatemic familiar tumoral calcinosis (*SAMD9*), in hepatitis B virus-associated hepatocellular carcinoma (*SAMD9L*), in MDS/AML with microdeletion in 7q21 (both genes), and in *SAMD9L*-haploinsufficient mice. Based on the observations in this murine model, the cytokine-receptor complexes cannot be properly degraded due to impaired endosomal function in *SAMD9L*-haploinsufficient cells, which results in constitutive intracellular signaling with prolonged cell survival(18). Moreover, *SAMD9L+/-* and *SAMD9L-/-* mice develop MDS with normo/hypercellular BM, drawing an a parallel to human *SAMD9L*-related MDS where the initial marrow hypocellularity associated with “toxic” mutation is restored upon the loss of mutant *SAMD9L* allele. This loss is accomplished through -7, del(7q) or UPD7q, and leads to proliferative advantage with clonal expansion. In our cohort, although initially all patients presented with
hypocellular marrow, longer observation periods in P1, P2, P4, and P7 revealed successive normalization of marrow cell content associated with increasing -7/del(7q) clone or appearance of UPD7q. In P1, the leukemic progression was further aggravated with the accumulation of typical MDS driver mutations in \textit{SETBP1}, \textit{KRAS} and \textit{EZH2} within the -7 clone. P5 also demonstrated an acquired splice site mutation in \textit{RUNX1}, a known leukemic driver gene. Notably, typical adult MDS driver mutations (i.e. \textit{TET2}, \textit{DNMT3A}, \textit{iDH1/2}) were not encountered in our cohort. This is in line with previously published findings discussing rather \textit{SETBP1}, RAS pathway mutations and \textit{RUNX1} (identified in our \textit{SAMD9L}-mutated patients) as recurrent drivers of pediatric MDS (10). Building on our observations we propose the following mechanism of MDS evolution in \textit{SAMD9L} disease: BM attempts to circumvent the toxicity of constitutional \textit{SAMD9L} mutation and selects for fitter, yet premalignant -7/del(7q) clones (with only one wildtype \textit{SAMD9L} copy), or benign clones with truncated \textit{SAMD9L} (Figure 3). Over time the resulting haploinsufficiency of tumor suppressor genes on 7q (e.g. \textit{EZH2} or \textit{CUX1}) in all patients likely provide the first step towards progression. Finally, additional driver somatic mutations might be encountered in some but not all patients.

Somatic revertant mosaicism has been reported in IBMFS with hypocellular BM, including telomeropathy with germline mutations in \textit{TERT}(35), and Fanconi anemia where mosaicism in blood occurs at rates of up to 140x higher than in general population(36, 37). However, in general, revertant mosaicism is a rare facet to clonal hematopoiesis because spontaneous correction of the pathogenic allele is a random event. In our study, we report two patients (P2 and P7) who presented with RCC/-7 at young age and demonstrated complete hematopoietic remission with normal cytogenetics throughout an observation period ranging from 16 to 20
years. The clinical picture of these patients fits the previously described transient monosomy 7 syndrome; to our knowledge 8 patients with primary MDS with transient -7 or del(7q) were reported in the literature (14-16, 38-40). Their age at diagnosis ranged from 8 months to 3 years, and spontaneous remission was achieved within 1-20 months from diagnosis. It seems that the -7 clones in our patients either spontaneously vanished or were outcompeted by “fitter” UPD7q-corrected clones with diploid copy of wild type \textit{SAMD9L} allele. Based on these observations, a watch-and-wait strategy might be proposed for younger patients with RCC/-7 who have no additional somatic driver mutations and are clinically stable. However, prolonged “watchful waiting” poses the risk of progression as witnessed in P1, who advanced to CMML and acquired oncogenic mutations 3.6 years after he was diagnosed of RCC.

In conclusion, our observations establish the molecular basis of a distinct subtype of familial MDS and point to the notion that MDS with chromosome 7 loss can be the sole and common manifestation of \textit{SAMD9L}-related disease. The negative mutational effect leads to escape and outgrowth of clones carrying -7/del(7q) with only wildtype \textit{SAMD9L} allele, which might spontaneously disappear or persist and provide the first step towards disease progression. Finally, this is the first description of long-term revertant mosaicism due to nonrandom UPD7q in \textit{SAMD9L} disease, and a plausible explanation for transient monosomy 7 syndrome.
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Provided in the supplement.

AUTHOR CONTRIBUTION

MWW, VBP, SS designed the study and wrote the manuscript. VBP, SS, DL, MV, YTB, MB, JH, HB, and MWW performed experiments and analyzed data. CMN, JCA, BS, and ME contributed to study conception and data interpretation. JB, GCS, ES, BS, GE, MN, BSch, IB, AS, UT, LN, HH, CMN were involved in patient care, testing procedures and sample acquisition. All authors contributed to writing and approved the manuscript.

CONFLICT OF INTEREST

The authors declare no conflict of interest.
REFERENCES


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Table 1. Clinical data of SAMD9L-mutated patients.

<table>
<thead>
<tr>
<th>Patient # (UPN); sex</th>
<th>Gestational age; measurements</th>
<th>Dysmorphic feat., neural symptoms</th>
<th>Prior medical problems</th>
<th>Timepoint; MDS subtype</th>
<th>PB findings (Plt, WBC; ANC x 10^9/L; Hb g/dL)</th>
<th>BM findings</th>
<th>Genotype notes</th>
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| P1 (D098); male     | 39w; 3,550g (P50-70); 55cm (PIQ) | none                             | recurrent RTI and endogenous eczeema since infancy, recurrent pneumonia at 6 mo, Plt↓, ANC↓ at 2 yrs; self-limiting seizures at 3 yrs | 3.4y rs; RCC | (
P1: L, WBC↓; ANC↓; MCV↓; Hb↓; HbF↑) | +++ vacuolat ion in M | 45,XY,-7 [8] / 46,XY [10] | 60% |
| P2 (D508); female   | 34w; 2,670g (P75-90); 49cm (PIQ) | none                             | recurrent RTI since infancy; endogenous eczeema since infancy | 2.0y rs; RCC | (Plt↓) | +++ vacuolat ion in M | 45,XX-7 [3] / 46,XX [18] | 16% |
| P3 (US1); female    | 40w; 4,080g (P90-97); 53cm (PIQ) | none                             | endogenous eczeema | 12mo; RCC | (Plt↓) | +++ vacuolat ion in M | 45,XX-7 [3] / 46,XX [18] | 16% |
| P4 (US2); male      | 41w; 3,540g (P75-95); 52cm (PIQ) | none                             | recurrent RTI and endogenous eczeema since infancy | 15mo | (Plt↓) | ++ | 46,XY [10] | normal |
| P5 (D637); male     | term: normal                  | none                             | panocytopenia and hypocellular marrow at 1.7 yrs | 7.7y rs; RCC | (Plt↓) | ++ | 45,XY,-7; del(18)q (q10q12) [4] / 46,XY [10] | 19% |
| P6 (D637); term: normal | none             | none                             | normal               | 42y rs; RCM 0 | (Plt↓) | +++ | 45,XY; del(15)(q10?q11) / 46,XY [10] | n.p. |
| P7 (SCB-4); female  | term: normal                  | panocytopenia and hypocellular marrow at 1.7 yrs | 2.1y rs; RCC | (Plt↓) | ++ | 45,XX-7 [4] / 46,XX [17] | -7 confirmed normal | 7% |

UPN, unique patient number; synd., syndrome; w, gestational week; RTI, respiratory tract infection (including cith, bronchitis, pneumonia); yrs, years of age; mo, months of age; Dx, diagnosis; RCC, refractory cytopenia of childhood; CMML, chronic myelomonocytic leukemia; BM, bone marrow; E, erythropoiesis, M, myelopoiesis; n.p., not performed; Plt, platelets; MCV, mean corpuscular volume (according to age); WBC, white blood cell count; ANC, absolute neutrophil count; Dysplasia: +, mild; ++, moderate; ++++, severe. N, normal.

*with occasional small dysplastic megakaryocytes. †51% of metaphases with monosomy 7, of those 17% additionally showed hypoploid metaphases with involvement of chromosome 9, 14, 19, 21.
### Table 2. Overview of germline and somatic mutations.

<table>
<thead>
<tr>
<th>Mut type</th>
<th>Patient # (UPN)</th>
<th>Time point, age</th>
<th>Germline status</th>
<th>Myeloid sample</th>
<th>Germline source</th>
<th>Conservation</th>
<th>Effect (SIFT/ExAC Conservation)</th>
<th>CADD Score</th>
<th>InAC burnout rate/% in population</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FAMILY I</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P1 (D084)</td>
<td>7yrs (K M M1)</td>
<td>60% / [6/16]</td>
<td>SAMD9L</td>
<td>c.4534G&gt; A</td>
<td>p.V1512M</td>
<td>BM-GR</td>
<td>High / Small</td>
<td>D/P/D/D(65%)</td>
<td>25.2</td>
</tr>
<tr>
<td>Father</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother</td>
<td></td>
<td></td>
<td>-</td>
<td>SAMD9L</td>
<td>c.4534G&gt; A</td>
<td>p.V1512M</td>
<td>PB</td>
<td>High / small</td>
<td>D/P/D/D(65%)</td>
</tr>
<tr>
<td><strong>FAMILY II</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P3 (US1)</td>
<td>20mo</td>
<td>16% / [0/16]</td>
<td>SAMD9L</td>
<td>c.563A&gt;G</td>
<td>p.V1512M</td>
<td>BM-GR</td>
<td>High / Small</td>
<td>D/P/D/D(65%)</td>
<td>25.2</td>
</tr>
<tr>
<td>Father</td>
<td></td>
<td></td>
<td>-</td>
<td>SAMD9L</td>
<td>c.563A&gt;G</td>
<td>p.V1512M</td>
<td>PB</td>
<td>High / small</td>
<td>D/P/D/D(65%)</td>
</tr>
<tr>
<td><strong>FAMILY III</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P5 (US7)</td>
<td>42yrs</td>
<td>- / [0/20]</td>
<td>PTEN</td>
<td>p.460P &gt; T</td>
<td>p.H880Q</td>
<td>BM-GR</td>
<td>Weak / Small</td>
<td>D/-/D/-(-)</td>
<td>17.9</td>
</tr>
<tr>
<td>Father</td>
<td></td>
<td></td>
<td>-</td>
<td>PTEN</td>
<td>p.460P &gt; T</td>
<td>p.H880Q</td>
<td>PB</td>
<td>Weak / small</td>
<td>D/-/D/-(-)</td>
</tr>
<tr>
<td><strong>FAMILY IV</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P7 (SC054)</td>
<td>2.3yrs</td>
<td>- / [0/25]</td>
<td>SAMD9L</td>
<td>c.3562C&gt;T</td>
<td>p.R1188X</td>
<td>BM-GR</td>
<td>High / Large</td>
<td>D/N/B/D(87%)</td>
<td>23.7</td>
</tr>
<tr>
<td>P1 (D084)</td>
<td>7yrs (K M M1)</td>
<td>60% / [6/16]</td>
<td>RUNX1</td>
<td>c.2602G&gt; A</td>
<td>p.D868N</td>
<td>BM-GR</td>
<td>High / Large</td>
<td>D/N/B/D(87%)</td>
<td>23.7</td>
</tr>
</tbody>
</table>

**Abbreviations:** Mut, mutation; UPN, unique patient number; BM, bone marrow; PB, peripheral blood; GR, granulocytes; HR, hair follicles; FB, fibroblasts; m., maternal origin; p., paternal origin; VAF, variant allele frequency; WES, whole exome sequencing; DS, targeted deep sequencing; Sanger, identified by Sanger sequencing; n. a., not available; yrs/mo, years/months after diagnosis. Evolutionary conservation scores: Phylop and PhastCons; PhysChem diff, physicochemical difference between amino acids. In-silico prediction: SIFT: T-tolerated, D-damaging; Mutation Taster: Disease causing, N-neutral; PolyPhen2: D-probably damaging, B-benign; PredictSNP consensus classifier: D-deleterious, N-neutral (% accuracy). Combined annotation-dependent depletion (CADD-score) of 20 means that a variant is among the top 1% of deleterious variants in the human genome. CADD-20 = 1%, CADD-30 = 0.1%, CADD-40 = 0.01%, CADD-50 = 0.001%. *mutation classified as acquired based on low allelic frequency.

**Gene annotations:** SAMD9L | NM_001303550.1, EZH2 | NM_152998, SETBP1 | NM_015539, KRAS | NM_004985.5, FANCM | NM_001308134, JAK3 | NM_002015, PTEN | NM_000314.4, RUNX1 | NM_00101890.0
FIGURE LEGENDS

Figure 1. Germline SAMD9L mutations in pedigrees with familial MDS.

(A) Identification of 4 pedigrees with MDS and monosomy 7 harboring germline heterozygous SAMD9L mutations: p.V1512M (pedigree I), p.R986H (pedigree II) and p.R986C (pedigree III), p.H880Q (pedigree IV), and somatic mutations: p.R1188X (P1) and p.S1317RfsX21 (P7). Dotted symbols indicate healthy mutation carriers. Sanger sequencing on DNA extracted from hair follicles (HR) confirmed germline status of mutations as visualized in electropherograms. Sequencing in P1 was performed on peripheral blood granulocytes (GR) revealing a minor mutational peak, corresponding to a variant allelic frequency of 8.3% by WES. In pedigree III, the mutation in P5 was confirmed in fibroblast (FB) DNA, while for P6 and remaining family members whole blood (WB) was analyzed. In pedigree IV other family members were not tested (n.t.). (B) SAMD9L and SAMD9 gene orientation on 7q22 in reverse strand direction (3’-5’). SAMD9L protein is coded by one exon and contains two known functional sites: N-terminal sterile alpha motif (SAM) and nuclear localization sequence (NLS). Four germline and two somatic (*) mutations were identified in SAMD9L. Germline missense mutations are evolutionarily highly conserved. (C) TA cloning of double mutated SAMD9L region of P1 revealed cis-configuration of mutations p.V1512M (germline) and p.R1188X (somatic) in 10 out of 172 clones tested.

Figure 2. Bone marrow findings in P1, P2 at different timepoint of disease course.

H&E staining of bone marrow (BM) at diagnosis of RCC in P1 showing dysplastic granulopoiesis with hypergranulation and a pseudo Pelger cell (top left), myeloblast and dysplastic eosinophil (top right). BM at diagnosis in P2 (synchronous with monosomy 7) showing hypergranulation and vacuolization in myelocytes, and dysplastic erythropoiesis with double nuclei (bottom left). Normal BM morphology in P2, 15 years after initial BM confirming spontaneous phenotype reversion (bottom right).

Figure 3. Mechanisms of clonal escape from SAMD9L germline mutations.

Multiple mechanisms of clonal escape from damaging germline missense SAMD9L mutations are observed and lead to complete (monosomy 7) or partial (deletion 7q) loss of chromosome 7 with decreasing mutant SAMD9L allele (red circles), both situations can lead to MDS development; UPD7q and truncating somatic SAMD9L mutations (green circles), which show benign outcome and contribute to normal hematopoiesis. Multiple clonal outcomes can occur in single patient.

Figure 4. Loss of mutated SAMD9L allele due to genomic deletion or mitotic recombination.
Variant allelic frequency (VAF) scores for chromosome 7 in P1 and P2. SNPs and Indels detected using whole exome sequencing (~4000 variants with a VAF score > 5% and < 95%), depict a complete loss of chromosome 7 in P1, as the VAF scores are either low or high. P2, unlike P1 demonstrates a partial loss of the Chr 7 after position 7q11.22 towards the q terminal site. The read depth of the SNPs for P2 was maintained throughout for Chr7 with no loss thus confirming the LOH is due to UPD and not -7q. Whole exome sequencing VAF values are marked by a star within the graph. Abbreviations: VAF, variant allelic frequency; UPD, uniparental isodisomy.

For P7, targeted NGS identified 14 informative (heterozygous) polymorphisms located on chromosome 7q with an average depth of 1036 reads (supplementary table 1). SNPs are represented in a VAF graph depicting the skewing of heterozygosity towards one allele occurring after position g.66098482 (rs3764903).

**Figure 5. Clonal evolution and spontaneous reversion due to UPD7q.**

Clonal evolution model in P2 (D154) depicting disease history during an observation period of 20 years. At diagnosis, initial bone marrow harbored monosomy 7 (77% by FISH and 51% by metaphase karyotyping). Blood counts normalized 3.7 years later and since then P2 maintained normal CBC until last follow-up at age of 22 years. From age of 12 years, repetitive yearly BM examinations revealed normocellular hematopoiesis with no dysplasia and normal cytogenetics. BM collected at age of 17 years (*) was subjected to whole exome and targeted deep sequencing. Germline heterozygous SAMD9L mutation p.V1512M was detected at a variant allelic frequency (VAF) of ~20%, corresponding to a clonal size of ~40% in a diploid chromosome 7 background. Concurrently, a spontaneous genetic correction of the SAMD9L locus occurred resulting from uniparental isodisomy (UPD)7q of paternal origin. This self-corrected clone occurred either initially (dotted line) or after termination of monosomy 7 and contributed to normal hematopoiesis. Abbreviations: Dx, diagnosis; pat, paternal origin; mat, maternal origin; UPD; uniparental isodisomy, LFU; last follow-up.

**Figure 6. Functional evaluation of SAMD9L mutations.**

(A-B) The effect of SAMD9L mutations on cell proliferation was assessed by dye dilution assays. 293FT cells were transiently transfected with TFP-SAMD9L wild type (WT), the disease-associated mutations p.R986C and p.V1512M, and the protective variant p.T233N previously reported by Tesi et al[20]. (A) Histograms depict the dye levels in transfected cells. Dye levels were monitored in TFP-transfected cells (filled grey histograms) and compared to cells expressing uniformly intermediate levels of TFP-SAMD9L wild type (blue histograms) or variants (red/orange lines), as indicated. A single representative experiment is shown. (B) Cumulative summary of three independent experiments on inhibition of cell proliferation associated with indicated TFP-SAMD9L mutations. Values (mean +/- SD) are calculated based on a scale defined by 0 (dye levels in TFP-transfected cells) and ~1 (dye levels in cells transfected with TFP-SAMD9L wild type). Unpaired t-test, two tailed: * p<0.05; ** p<0.005; *** p<0.001.
Figure 1. Germline SAMD9L mutations in pedigrees with familial MDS

A

I. c.4534G>A, p.V1512M

II. c.2957G>A, p.R986H

III. c.2956C>T, p.R986C

IV. c.2640C>A, p.H880Q

B

Chr:7q21
GR.Ch37/hg19

g.92,765,304 - 92,760,530

SAMD9L

SAMD9

NH2

SAM

NLS

p.H880Q

p.R1188X

p.R1188X

p.R1512M

p.R1512M

C

7q21

1. WT
2. p.V1512M

No. of colonies

0

50

100

150

170

153
Figure 2. Bone marrow morphology in P1 and P2

P1 initial

P1 initial

P2 initial

P2 at remission (+15 years)
Figure 3. Mechanisms of clonal escape from SAMD9L germline mutations

**GERMLINE** (polyclonal hematopoiesis)

**heterozygous**

\[ \text{germline mutation} \]
\[ \text{somatic mutation} \]

\[ \text{WT} = \text{mut} \]
\[ 50:50\% \]

**MECHANISMS OF CLONAL ESCAPE**

- **Monosomy 7**:
  - \[ g^{\text{SAMD9L mut}} \downarrow \]
  - MDS
  - Disappearing -7
  - **P1, P3-6**

- **Deletion 7q**:
  - \[ g^{\text{SAMD9L mut}} \downarrow \]
  - MDS
  - **P4**

- **UPD7q**:
  - \[ g^{\text{SAMD9L mut}} \downarrow \]
  - Clonal expansion & sustained self-correction
  - **P2, P7**

- **Truncating**
  - \[ g^{\text{SAMD9L mut}} \downarrow \]
  - Rescue by truncating SAMD9L mutation (insufficient for complete correction)
  - **P1, P7**
Figure 4. Loss of mutated SAMD9L allele due to genomic deletion or copy-number neutral recombination.

SAMD9L (7q21)

P1

VAF (%)

100
80
60
40
20
0

p.V1512M (VAF)

UPD 7q11.22-ter

21.5%

P2

VAF (%)

100
80
60
40
20
0

p.H880Q (VAF)

26%

P7
Figure 5. Clonal evolution and spontaneous reversion due to UPD7q

Monosomy 7
SAMD9L WT
~77%

UPD 7q11.2-qter
SAMD9L WT
~60%

Native hematopoiesis
SAMD9L p.V1512M ~40%

<table>
<thead>
<tr>
<th>P2: age, years</th>
<th>Normal CBC</th>
<th>Normal Cytogenetics</th>
</tr>
</thead>
<tbody>
<tr>
<td>45,XX, -7</td>
<td></td>
<td></td>
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<tr>
<td>2 (Dx)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.7</td>
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<td>14</td>
<td></td>
<td></td>
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<tr>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18</td>
<td></td>
<td></td>
</tr>
<tr>
<td>22 (LFU)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 6. Functional evaluation of SAMD9L mutations.

A

WT

p.T233N

p.R986C

p.V1512M

B

proliferation index


TFP-0SAMD9L
Figure S1. Shared variants found by WES in P1 and P2

Legend: The Y axis corresponds to the average VAF % of all detected shared mutations in both siblings and the X axis the calculated CADD-scores. The vast majority of mutations show a low degree of deleteriousness with a CADD-score lower than 20. Framed in red, 50 highly deleterious variants of which 48 were exonic. Among them, germline mutations in *SAMD9L* (p.V1512M) and *PTEN* (p.Y188C) were observed.
Figure S2. Individual mutations found in P1 by WES

Legend: Individual exonic variants in P1 and P2. Highly deleterious (CADD>20) variants were 58 and 45 variant in P1 and P2 respectively. Known leukemic driver genes were affected uniquely in P1 in addition to a novel non-sense variant in the SAMD9L gene. In contrast, no significant mutations involved in blood disease or cancer were found in P2.
Table S1. Allelic frequencies of informative chromosome 7q polymorphisms in P7

<table>
<thead>
<tr>
<th>Gene</th>
<th>Isoform</th>
<th>Genomic Position</th>
<th>SNP</th>
<th>cDNA</th>
<th>VAF (allele 1)</th>
<th>VAF (allele 2)</th>
<th>dbSNP:MAF</th>
</tr>
</thead>
<tbody>
<tr>
<td>KCTD7</td>
<td>NM_001167961</td>
<td>66098482</td>
<td>rs3764903</td>
<td>c.314+51G&gt;A</td>
<td>Het. G</td>
<td>52%</td>
<td>A</td>
</tr>
<tr>
<td>GTPBP10</td>
<td>NM_033107</td>
<td>89984409</td>
<td>rs42664</td>
<td>c.329A&gt;G</td>
<td>Het. A</td>
<td>28%</td>
<td>G</td>
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<tr>
<td>DYN1I1</td>
<td>NM_004411</td>
<td>95709602</td>
<td>rs42083</td>
<td>c.1702-73C&gt;T</td>
<td>Het. C</td>
<td>24%</td>
<td>T</td>
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<td>DYN1I1</td>
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<td>150768786</td>
<td>rs2303937</td>
<td>c.2160G&gt;A</td>
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<td>PRKAG2</td>
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<td>151504499</td>
<td>rs10257529</td>
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<td>Het. C</td>
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<td>Het. A</td>
<td>25%</td>
<td>G</td>
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</table>

Average read depth obtained for all 14 polymorphisms: 1036x.
CASE DESCRIPTIONS

In **family I**, the older brother P1 suffered from recurrent respiratory tract infections (RTI) and endogenous eczema manifesting in infancy. The first documented complete blood count (CBC) during an infectious episode at the age of 6 months showed pancytopenia (platelets $39 \times 10^9$/L, neutrophils $0.67 \times 10^9$/L, Hb 7.7g/dL). Anemia spontaneously improved but platelet and neutrophil counts remained low normal. The first bone marrow (BM) aspirate performed at the age of 2 years showed a reduced cell content and dysplasia with vacuolization in the myeloid and erythroid lineages and BM eosinophilia. Hemoglobin F, Vitamin B12, folate and immunoglobulin levels as well as lymphocyte subpopulations were within normal range and there were no signs of autoimmunity. The patient remained stable without transfusions. A second BM aspirate performed at the age of 3.4 years confirmed myelodysplasia (Figure 2); chromosomal studies showed monosomy 7 in 6 of 16 metaphases. P1 was diagnosed with refractory cytopenia of childhood (RCC). He suffered transient cerebral seizures without the need of anticonvulsant therapy, while a cranial CT scan was normal. At the age of 6.5 years, he presented with hemorrhagic varicella with secondary pneumonia and required platelet transfusions for the first time. At the age of 7 years, he had developed hepatosplenomegaly and showed 21% blasts on peripheral blood smear. There was no spontaneous growth in *in vitro* GM-CSF hypersensitivity assay. AML induction chemotherapy was complicated by bacterial sepsis and respiratory failure. Following recovery, thioguanine maintenance therapy was administered. He received an HSCT from a matched unrelated donor (MUD). Hematological reconstitution was slightly delayed (WBC day +19, platelets day +63, RBC day +49) with stable complete chimerism. At day +138 late, grade III acute GvHD (gut stage 3, liver stage 2, liver stage 2) manifested, and he died 10 days later from acute CNS hemorrhage.

P2 is the younger sister of P1 and presented with recurrent respiratory tract infections at the age of 18 months. The first documented CBC at the age of 2 years revealed thrombocytopenia (Table 1). BM was hypocellular with severe dysplasia, vacuolization similar to what had been observed in P1 (Figure 2). Monosomy 7 was detected in 77% FISH-interphases and 51% metaphases with hypoploidy metaphases in one subclone (Table 1). After the death of her brother, P2 did not visit the hematology clinic until two years later for tonsillectomy due to recurrent RTI. Her CBC revealed only mild thrombocytopenia ($85 \times 10^9$/L) and leukopenia ($4.4 \times 10^9$/L) without neutropenia. When she was 5.7 years old her CBC normalized and remained within the normal range since. A second BM examination at the age of 12 years revealed normal cellularity with slight dysplastic changes and normal cytogenetics. All subsequent marrow analyses in nearly yearly intervals were normal without any signs of MDS or monosomy 7 (Table 1, Figure 2). P2 did not suffer from infectious episodes and at last follow-up (FUP) at the age of 22 years she was in good general condition.

In **family II** the older sister (P3) had unremarkable postnatal development with the exception of chronic endogenous eczema, and petechial rash manifesting at the age of 1 year. No recurrent infections were observed. Her CBC performed during routine 18 months evaluation revealed thrombocytopenia, neutropenia and elevated MCV. Initial BM was hypocellular with mild erythroid dysplasia, and without increase of blasts. Cytogenetics uncovered monosomy 7 in 3/21 metaphases and 16% FISH-interphases. Six months later she was successfully transplanted from a MUD after myeloablative conditioning. At last FUP she was 6 years 4 months old and in good general condition.
P4 is the younger brother of P3. He suffered from recurrent RTI since infancy. His initial hematologic presentation during an episode of acute otitis media at the age of 12 months was pancytopenia with severely hypocellular BM and dysgranulopoiesis (Table 1). Evolving MDS-RCC or toxic stromal damage secondary to a viral insult was suspected. Viral workup was unremarkable. His CBC recovered and subsequent BM examinations demonstrated normocellular BM without significant dysplasia. However, 3 months after initial presentation, a small monosomy 7 clone was detected by FISH in 11/200 interphase-cells but was absent in metaphase cytogenetics. Finally, 3 months later, two independent clones emerged, with -7 and del7q in 6/20 and 4/20 metaphases, respectively (Table 1). At last FUP, P4 was 21 months old and undergoing MUD-HSCT.

In family III, the affected son (P5) and his father (P6) did not have any infectious or hematologic problems until they presented with pancytopenia and were diagnosed with hypocellular MDS at the age of 7.7 and 42 years, respectively. Complete or partial loss of chromosome 7 in addition to other concomitant lesions (Table 1) were found in BM. Both P5 and P6 were transplanted from an unrelated donor, or HLA-identical brother at the age of 8 and 43 years, respectively. They were alive at last FUP when they were 17.4 and 54 years old.

In family IV, P7 was the affected child of non-consanguineous parents with negative family history for hematopoietic or neurologic diseases. At the last FUP, P7’s father had been evaluated by neurologists for unclear ataxia symptoms (his SAMD9L status is unknown). The girl presented at the age of 1 year and 8 months with pancytopenia and petechial rash on the trunk. Initial bone marrow showed a cellularity of 15%. Marrow analysis performed 5 months later revealed normal cellularity and dysplasias compatible with the diagnosis of MDS-RCC, while cytogenetics showed complete monosomy 7 in 4 out of 21 metaphases as confirmed by interphase FISH. Unexpectedly, cytogenetics normalized 3 months later, cellularity gradually increased, and there were only slight dysplastic traits observed in the myeloid lineage. At that time point NGS analysis in BM specimen identified a large UPD7q clone (double wildtype for SAMD9L). Subsequent marrow examinations and CBC analyses were normal and the patient remained healthy until last follow-up at the age of 18 years.
SUPPLEMENTARY METHODS

Genomic studies
Genomic DNA from P1 and P2 was subjected to exploratory whole exome sequencing (WES). After exon region capture using Agilent SureSelect v5.0 (Agilent, Santa Clara, CA) and library construction by Illumina TruSeq (Illumina, San Diego, CA), 100bp paired-end sequencing was performed on Illumina HiSeq2000 instrument. Following dynamic trimming, the resulting reads were aligned to the hg19 reference genome with BWA20 followed by base quality score recalibration, indel realignment, duplicate removal, SNP and INDEL discovery using standard filtering parameters, and variant quality score recalibration according to GATK1,2. Confirmatory deep sequencing was performed for SAMD9L in P1 and P2 and P4 for respective mutations (Table 2). P1, P2, P5 and P7 were subjected to an in house 28-gene pediatric MDS panel (University of Freiburg, Germany). P4 was analyzed using a “57-genes myeloid malignancies” NGS panel (ARUP laboratories, Salt Lake City, USA) and a “86-genes bone marrow failure” NGS panel (Claritas Genomics, Cambridge, USA).

All mutations identified by NGS were validated using Sanger sequencing. For germline confirmation, DNA from skin fibroblast and/or hair follicles was extracted using automated extraction system (AS 1290 kit, MaxWell 16 System, Promega). The targets were amplified using Amplitaq Gold Polymerase (Lifetechnologies, USA) and sequenced as previously described4.

Bioinformatics
The variant annotation was compiled using ANNOVAR5 from the 1000 Genomes Project requiring a minimum of 8 reads and a variant allele frequency > 0.05%. SNPs reported at a minor allele frequency <0.1% in the ExAc Browser were considered rare. We applied a minimum quality score of 30 (Q30, 99.9% base call accuracy) to minimize the incorrect base calls (Q30=1/1000 incorrect base calls). A minimum coverage of 150X was employed for genetic variants with a variant allelic frequency (VAF) ≥ 5%. Those quality thresholds were recommended by the Illumina Technical Notes: 1) Quality Scores for Next-Generation Sequencing, 2011, 2) Somatic Variant Caller, 2012.

The analysis of identified variants was performed using Alamut Visual 2.9 (Interactive Biosoftware, Rouen, France) and evaluated with open source genomic databases i.e. catalogue of somatic mutations in cancer (COSMIC v77)6, ExAC Browser (Version 0.3)7, UniProtKB/Swiss-Prot database, Ensembl genome browser 85 and dbSNP - NCBI - National Institutes of Health. The degree of deleteriousness was calculated using a PHRED-like scoring system “Combined Annotation Dependent Depletion” (CADD) - score. Using this meta-annotation tool, CADD-scores are assigned to genetic variants using an algorithm that creates a consensus based on 60 functional prediction tools. A CADD score of 10 refers to the top 10% of deleterious variants detected in the human genome. CADD score above 20 represents the top 1% of deleterious variants.8 Finally, the pathogenicity was evaluated using standard in-silico predictors: PolyPhen29, SIFT10, Mutation Taster11 and PredictSNP12. The evolutionary conservation of nucleotides/amino acids across species and the physicochemical difference between amino acids was estimated by PhyloP, PhastCons and Grantham score respectively13. Sanger sequencing was performed using a Sanger ABI sequencer. Sanger sequences were visualized in Sequence Pilot (SeqPilot v 3.5.2) and CodonCode Aligner v6.0.2.
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SUPPLEMENTARY REFERENCES


