New KCNN4 Variants Associated With Anemia: Stomatocytosis Without Erythrocyte Dehydration

B. Allegrini, S. Jedele, L. David Nguyen, M. Mignotet, R. Rapetti-Mauss, C. Etchebest, O. Fenneteau, A. Loubat, A. Boutet, C. Thomas, J. Durin, A. Petit, C. Badens, L. Garçon, L. Da Costa, and H. Guizouarn

The K⁺ channel activated by the Ca²⁺, KCNN4, has been shown to contribute to red blood cell dehydration in the rare hereditary hemolytic anemia, the dehydrated hereditary stomatocytosis. We report two de novo mutations on KCNN4, V222L and H340N, characterized at the molecular, cellular and clinical levels. Whereas both mutations were shown to increase the calcium sensitivity of the K⁺ channel, leading to channel opening for lower calcium concentrations compared to WT KCNN4 channel, there was no obvious red blood cell dehydration in patients carrying one or the other mutation. The clinical phenotype was greatly different between carriers of the mutated gene ranging from severe anemia for one patient to a single episode of anemia for the other patient or no documented sign of anemia for the parents who also carried the mutation. These data compared to already published KCNN4 mutations question the role of KCNN4 gain-of-function mutations in hydration status and viability of red blood cells in bloodstream.

Keywords: Hereditary Xerocytosis, Stomatocytosis, red blood cell, Gardos, KCNN4

INTRODUCTION

The rare disease dehydrated hereditary stomatocytosis (DHSt or hereditary xerocytosis, HX) is an autosomal dominant hemolytic anemia, characterized by an alteration of the cation permeability of red blood cell (RBC) (Badens and Guizouarn, 2016; Gallagher, 2017; Flatt and Bruce, 2018). In 2012, the identification of PIEZO1 mutations in families suffering DHSt paved the way toward the genetic cause of this long-known hemolytic anemia (Zarychanski et al., 2012; Albuisson et al., 2013; Andolfo et al., 2013). Three years later, with the identification of mutations in KCNN4, another genetic cause of DHSt was claimed: “two genetic causes for a single RBC pathology” (Grygorczyk and Mohandas, 2015). KCNN4 is a Ca²⁺-activated K⁺ channel (also named SK4 or Gardos channel in RBC) (Gárdos et al., 1977; Maher and Kuchel, 2003) and PIEZO1 is a non-selective cation channel, permeable to Na⁺ and K⁺ and divalent cations Ca²⁺ and Mg²⁺ (Gnanasambandam et al., 2015) activated by mechanical stimuli (Coste et al., 2012). Numerous gain-of-function mutations in PIEZO1 were identified in families with DHSt (Picard et al., 2019; More et al., 2020) and some of them were shown to modify the channel gating leading to longer open state that should increase Ca²⁺ concentration in
RBC (Albuisson et al., 2013; Cahalan et al., 2015; Danielczok et al., 2017; Glogowska et al., 2017). This Ca$^{2+}$ concentration rise can activate KCNN4, which leads to RBC dehydration (Syeda et al., 2015). The large anion conductance of erythrocyte membrane allows the loss of K$^+$ when intracellular Ca$^{2+}$ increase opens KCNN4. This KC$^+$ loss is accompanied by osmotically linked water. The Ca$^{2+}$-induced K$^+$ loss, named the Gardos effect, is of primary importance in RBC physiology (Gárdos et al., 1977; Maher and Kuchel, 2003). These cells do not regulate their volume. Hence, the dehydration resulting from KCNN4 activation is irreversible and this is expected to alter the RBC rheological properties. In sickle cell disease, the Gardos effect contributes to RBC sickling, the S-S hemoglobin having a greater tendency to polymerize following dehydration than A-A hemoglobin (Brugnara et al., 1993; De Franceschi et al., 1994). This mechanism has also been proposed to participate in the senescence of RBC. In a recent study, Rogers and Lew suggested that successive micro-activations of KCNN4 slowly dehydrate the RBC contributing to its densification, a marker of senescence (Rogers and Lew, 2021). In the case of PIEZO1 gain-of-function mutations linked to DHSt, a chronic stimulation of KCNN4 is considered as the cause of the observed RBC dehydration. In patients’ RBC with different PIEZO1 mutations, KCNN4 activation appears as the sole effector of this dehydration (Rapetti-Mauss et al., 2017).

In patients’ RBC carrying KCNN4 mutations, the channel conductance was increased (Rapetti-Mauss et al., 2016; Rivera et al., 2017; Fermo et al., 2020). Moreover, the mutations changed channel gating mainly by modifying its Ca$^{2+}$ sensitivity (Garneau et al., 2009; Rapetti-Mauss et al., 2016).

Despite leading to a more active channel, the gain-of-function mutations in KCNN4 are not systematically linked to RBC dehydration, and routine hematological tests failed to clearly diagnose DHSt (Picard et al., 2021). Nonetheless, these KCNN4 mutations are associated with anemia that is often severe, especially in childhood and fetal life (Rapetti-Mauss et al., 2015).

Our present study was designed to reinforce our knowledge about KCNN4 mutations and DHSt. Two de novo KCNN4 mutations were identified in two unrelated families leading to the amino acid substitution V222L and H340N. This later had mutations were identified previous observation with the majority of the KCNN4 mutations, our data did not correlate KCNN4 gain-of-function mutations with RBC dehydration, raising the question of classifying this pathology as a DHSt. Moreover, it emphasized the difficulty to diagnose altered RBC permeability facing KCNN4 mutations and the great variability in RBC phenotypes associated with KCNN4 gain-of-function mutations.

**MATERIALS AND METHODS**

**Patients:** Patients had been referred to our Hematology Diagnostic laboratory (AP-HP) for phenotypic and genotypic explorations. The lab is quality certified for molecular screening analysis for “targeted-NGS red cell and erythropoiesis defects” (Cofrac Iso15189) and labeled “Reference Medical Biological Laboratory” for red cell membrane diagnosis (LBMR July 2021). The patients exhibited chronic hemolytic anemia/hemolysis signs or had been followed up for uncharacterized red blood cell membrane disorder.

**Red cell and reticulocyte indices, EMA test, and Ektacytometry:** All blood samples were collected on EDTA and shipped at 4°C after blood was drawn along with a blood smear. Samples should be delivered to our laboratory within 48h after blood collection. RBC indices including hemoglobin concentration, hematocrit, mean cell volume (MCV), mean corpuscular hemoglobin concentration (MCHC), mean hemoglobin content (MHC), RBC volume distribution (RDW), and reticulocyte count and distribution have been measured for each sample using a hematological analyzer (XN, Sysmex, Kobe, Japan). Blood smears stained with May Grünwald Giemsa (MGG) were carefully examined and blind diagnostics of RBC morphology abnormalities were validated independently by two cytologists prior to additional analysis. The EMA test has been performed according to the recommendation (Girodon et al., 2008; Da Costa et al., 2016) with modification (Da Costa et al., 2016). The mean fluorescence intensity (MFI) for each sample was compared to three age-matched controls collected on the same day. A ratio of the mean fluorescence for patient RBC to the mean fluorescence for the three controls was derived (mean of three age-matched control MFI–patient MFI/mean of three age-matched control MFI). Ektacytometry LoR Ra Max Sis (Mechatronics instruments BV®, Zwaag, Netherlands) has been performed as previously described (Mohandas and Chasis, 1993; Mohandas and Gallagher, 2008; Da Costa et al., 2016). Blood samples (minimum of 100 μl) were analyzed by ektacytometry freshly and in any case before 48h after blood sample collection. Briefly, samples were subjected to increasing shear stress and an osmotic gradient and the laser diffraction pattern through the RBC suspension were recorded. The RBC shape goes from circular to elliptical as shear stress increases. From these measurements, a deformability or elongation index for the cells can be derived. Three distinct features of the osmotic gradient ektacytometry profiles are the Omin, the DImax, and the O’ or hyperpoints. The Omin point corresponds to the osmolality at the minimal deformability in hypoosmolar area or at the osmolality when 50% of the RBC hemolyzed during the regular osmotic resistance test. It reflects the surface-to-area ratio. DImax corresponds to the maximal deformability index or elongation index (EI). The hyperpoint or O’ corresponds to the osmolality at half of the DImax and reflects the hydration state of the cells. Ektacytometry enables simultaneous analysis of three major RBC properties, RBC cell geometry, viscosity, and deformability, under the osmocan application of the ektacytometer.

**Genotype characterization:** Genomic DNA was extracted from blood lymphocytes. Written informed consent was obtained from affected individuals and/or parents prior to inclusion in this study, which was performed in accordance with the ethical standards of the Declaration of Helsinki. The targeted New Generation Sequencing (t-NGS), developed in the
lab, is a Roche “NimbleGen SeqCap EZ” library and an illumina flowcell (Flowcell standard 2*150) with a library of 187 genes including 93 genes for red cell disorders and erythropoiesis defects. Among them 21 genes were for red cell membrane disorders. The sequences have been run on a Miseq or a Nextseq in a genetic platform (Pr A. Verloes, Genetic Department, R. Debré hospital, Paris). Sequences have been analyzed on CLC Biomedical Work Bench and allelic variations have been interpreted with Qiagen Clinical Insight (QCI), Alamut visual (Sophia Genetics), and Varsome (Kopanos et al., 2019; Baux et al., 2021).

**Red cell cation content and volume measurements:** Fresh venous blood was obtained by venipuncture in EDTA collecting tubes from informed patients and healthy volunteers. Blood samples were received in the laboratory within 24–48 h traveling at room temperature. For vanadate experiments: blood was washed four times at room temperature in a medium containing (in mM): NaCl (145), KCl (5), MgSO4 (2), CaCl2 (1), and Hepes/NaOH, pH 7.4 (10). Red cell suspension was then incubated at room-temperature 25% hematocrit with 0.5 mM ouabain, and 5 mM vanadate was added alone or with 4 mM Senicapoc. A few minutes before sampling time, 400 µl of cell suspension was taken to three nylon tubes that were centrifuged for 10 min at 4°C, 20 000 g at the exact sampling time. The pellet of red cells was extracted and immediately weighted. Then, dry weight was measured after overnight heating (80°C). Water content was calculated with a correction of 3.64% corresponding to the trapped medium between packed cells. Intracellular ions were extracted from dried pellets by overnight incubation at 4°C in 5 ml milliRho water (Millipore). Two percent perchloric acid was then added to precipitate proteins, samples were centrifuged for 10 min 20 000 g, and supernatant was collected for Na+ and K+ quantification by flame spectroscopy with an Eppendorf ELEX6361.

**Intracellular Ca2+ measurement:** RBCs were washed two times (800 g, 5 min, and 4°C) in Ringer without Ca2+ to remove buoyy coat by aspiration. Four microliters of packed RBC were loaded with 2.5 µl of 1 mM Fluo-4 AM stock solution in 500 µl Ringer without Ca2+, 37°C, and 30 min. The Fluo4-loaded RBC suspension was directly used to quantify intracellular Ca2+ concentration (25 µl of RBC suspension in 975 µl Ringer without Ca2+ in FACs tubes) by measuring fluorescence with a FACs Fortessa BD. Internal RBC fluorescence was assessed on RBC treated without Fluo-4 AM. For vanadate experiments, the Fluo4-loaded RBC suspension was diluted 40 times in Ringer with 1 mM Ca2+ and 5 mM vanadate at time 0 and intracellular fluorescence was measured with FACs Fortessa BD as a function of time.

**HEK293T cells transfection:** HEK293T cells were grown in DMEM glutamax (Gibco) 10% FBS penicillin–streptomycin. Cells were co-transfected with WT or point-mutated pcDNA3-KCNN4-HA and pRES-eGFP (ratio 10:1) using CaPO4, WT pcDNA3-KCNN4-HA was a kind gift of Len Kaczmarek laboratory. Sixteen hours later, cells were washed twice with PBS, and patch clamp was done on fluorescently labeled cells. Point mutations were done by PCR on pcDNA3-KCNN4-HA with the proofreading DNA polymerase pfu-Turbo and primers covering 16 nucleotides upstream and downstream the single point mutation C1018 A for H340N or G649C for V222L. The pcDNA3-KCNN4-HA H340N or V222L clones used in the study were sequenced entirely.

**Protein expression assay:** HEK293T cells were grown to 70% confluence in DMEM glutamx 10% FBS penicillin–streptomycin in 60 mm Petri dishes (Starsted) and transfected with CaPO4 with 5 µg of DNA: pcDNA3-KCNN4-HA WT, V222L, or H340N per 5 ml cell culture. Transfected medium was removed after 6h. After 24h of expression, cells were biotinylated following manufacturer instructions (Pierce cell surface protein assay), and then lysed. The lysate was loaded on avidin-agarose beads (Pierce cell surface protein assay) to isolate the biotinylated fraction. Total fraction and biotinylated fraction were subjected to SDS-PAGE western blot. Migration of proteins was made at 120 V during 90 min in 10% acrylamide gel. Proteins were then transferred to PVDF using wet transfer protocol for 1h at 100 V and blocked in blocking solution (BS: 5% low-fat milk in TBS-tween 0.1%). Immunolabelling was done using primary antibodies: anti-KCNN4 (Proteintech, Rabbit, 1: 1,000), anti-Ecadherin (Cell Signaling, Mouse, 1:5,000), and anti-GAPDH (Calbiochem, Mouse, 1:200,000) for 1 h 30 min at room temperature in BS and HRP coupled secondary antibodies for 50 min at RT using: anti-rabbit (1:2,000, DAKO) and anti-mouse (1:5,000, DAKO). HRP-labeled proteins were revealed with Enhanced Chemiluminescent solution (Millipore) with a Fusion FX EDGE.

**Patch-clamp electrophysiology:** Glass pipettes (Brand, Wertheim, Germany) with final resistance of 3–5 MΩ were made on a horizontal pipette puller (P-97, Sutter Instrument, Navato, CA). All patch-clamp experiments were performed with a PC-controlled EPC9 patch-clamp amplifier (HEKA, Lambrecht/Pfalz, Germany). Currents were acquired and analyzed with Pulse and Pulsefit software (HEKA).

Currents were measured in whole-cell configuration with bath solution (mM): NaCl (145), CaCl2 (2), KCl (5), MgCl2 (1), and HEPEs (10), pH 7.4 adjusted with NaOH. Pipette solution (mM): KCl (145), MgCl2 (1), HEPEs (1), and EGTA (1), pH 7.2 adjusted with KOH. Free Ca2+ concentrations were adjusted by adding CaCl2 using Ca-EGTA Calculator v1.3. Currents were measured at room temperature using a ramp protocol from -120 to +80 mV from a holding potential of -60 mV (sampling frequency 10 kHz; filtered 5 kHz).

**KCNN4/CaM complex modeling:** Three full-length models of the KCNN4/CaM complex were constructed by assembling KCNN4 C-ter helices (residues 376 to 415, PDB ID: 6CNM) with the main part of the channel in three distinct conformational states (inactivated, activated/closed, and activated/open) with CaM and Ca2+ ions (PDB IDs: 6CNN, 6CNN, and 6CNO, respectively) (Lee and MacKinnon, 2018). Missing CaM N-lobe in the first structure was modeled by adding the N-lobe of the second structure and changing some torsion angles in CaM linker region, so that the CaM N-lobe is kept in solvent as described in the associated publication. Missing loops and residues were modeled using MODELLER (Webb and Sali, 2016) and residue protonation states were predicted at pH 7 using PROPKA.
Models of V222L and H340N KCNN4 mutants were obtained by replacing the side-chain of the mutated residue. Resulting structures were embedded in pure symmetric POPC membranes with 150 mM of KCl using CHARMM-GUI Membrane Builder server (Wu et al., 2014) and further relaxed using 1 µs molecular dynamics simulations carried out with GROMACS (Abraham et al., 2015). Models were represented using PyMol (Shrödinger L., DeLano W, 2020).

Residue conservation: Residue conservation analysis was based on the comparison of 3944 sequences collected from the NCBI non-redundant protein sequence database using blastp tool with human KCNN4 sequence (UniProt ID: O15554) as input query. Only sequences with sizes lower than 900 residues or with query coverage values higher than 66% were kept. Synthetic constructs were removed. Multiple alignments and associated weblogs were obtained using the EBI clustal omega tool (Sievers et al., 2011) and Weblogo server (Crooks et al., 2004). Conservation frequency of the KCNN4 residues was calculated as the occurrence ratio of the KCNN4 residue at the given position in the multiple sequence alignment.

RESULTS

Clinical Description of the Cases

Two subjects from two unrelated families were enrolled in this study after informed consent. The proband A had history of severe anemia requiring transfusion every 5–6 weeks since the age of 2 months when anemia was diagnosed. There was no splenomegaly, iron chelation was started at the age of 3 years. Blood withdrawal for permeability measurements was done on two occasions 3 months after the last transfusion.

The proband B was born after an uneventful pregnancy from non-consanguineous parents and delivery at 40 weeks by caesarean operation. At 9 months in a routine examination, he presented icterus and pallor requiring hematologic investigation in hospital. Results are presented in Table 1. Hemoglobin electrophoresis was normal (HbF 3.3%), G6PD and PK were normal, with no sign of infection, and Coombs test was negative. Bone marrow examination showed erythroid hyperplasia with discrete dyserythropoiesis. He was transfused and a steroid therapy had been started for 3 months. At the end of the steroid therapy, hemoglobin was stabilized. Growth and psychomotor development were normal and 4 years later, hematological parameters were normal (Table 1).

Genetics

Two allelic variations have been identified in KCNN4 gene (NM_002250.2) in two unrelated probands: 1) proband A carried a heterozygous missense allelic variation in exon 3 of KCNN4: c.664G > C; p.(Val222Leu) (Figure 1A), variant without significance [VUS or Class 3 (Richards et al., 2015; Amendola et al., 2016)], sift damaging. Mutation Taster disease causing, absent from the database. The variation has been inherited from the mother. 2) proband B carried a heterozygous missense allelic variation in exon 6 of KCNN4: c.1018C > T; p.(His340Asn)
(rs76935412) (Figure 1B), likely pathogenic (class 4, Varsome in silico prediction) variant, sift damaging, Mutation Taster disease causing, level pathogenic, MetaLR damaging, allele frequencies of 0.16%. The variation has been inherited from the mother.

**Hematological Parameters**

Table 1 compared hematological data coming from the new cases with H340N or V222L KCNN4 mutations with previously described RBC from patients carrying the R352H or V282M KCNN4 mutations. Anemia is a common feature of the different cases. This anemia is compensated in the family with V282M mutation showing a normal level of hemoglobin associated with hyper-reticulocytosis. The other cases exhibited a severe anemia not compensated by high reticulocytosis. The mutation H340N KCNN4 was correlated with a single episode of anemia which had not been further observed. In contrast, proband A with V222L KCNN4 mutation exhibited a severe uncompensated anemia associated with major dyserythropoiesis that could not be explained genetically, but a defect in iron/heme metabolism has been suggested.

MCHC is in the normal range or only few deciles above maximum. However, the MCHC has been reported at 37.6 g/dL in proband B at the time ektacytometry was performed (1.3 years) away from the hemolytic crisis (9 months). The size of RBC is most of the time larger than control (macrocytosis), while normal in probands A and B. These observations might result from the hyper-reticulocytosis: the maximum MCV is increased by 6–18% which could result from a 5–17% reticulocytes.

**Figure 1** | Blood smear, ektacytometry and genetics. DNA sequencing on proband A (A) and proband B (B). (C): Blood smear on proband A with her mother (mutation KCNN4 V222L). (D): Ektacytometry on proband A with her mother and control. (E): blood smear done on proband B RBC (mutation KCNN4 H340N) and his father and mother. Ektacytometry on proband B (F) and proband B’s mother’s and father’s RBC (G).
Blood smear showed anisopoikilocytosis (Figure 1C), with rare spheroctyes (purple arrows), rare elliptocytes (blue arrows), rare stomatocytes (black arrows), few RBC fragments (yellow arrows), and erythroblastemia for proband A with V222L KCNN4 mutation. Proband A’s mother’s blood smear did not show noticeable erythrocyte anomaly, except rare elliptocytes and very rare stomatocytes (Figure 1C). The ektacytometry indicated altered RBC deformability for proband A without any shift in osmotic resistance (Figure 1D). Omin point was normal while hyperpoint was shifted to the right. Proband A’s mother’s RBC deformability was normal and hyperpoint exhibited the same shift to the right as her daughter. For proband B, blood smear strikingly exhibited few stomatocytes [black and red (smile feature) arrows] (Figure 1E). The ektacytometry indicated normal deformability with a shift to the left of the hyperpoint in accordance with the increased MCHC at 37.6 g/dL in proband B at the time of the study (Figure 1F). The mother and the father exhibited normal ektacytometry curves (Figure 1G) and the mother, who carried the same allelic variation as his son, exhibited only rare stomatocytes, rare target cells (blue arrows), and acanthocytes (green arrows), while the father exhibited only few target cells (blue arrows) (Figure 1E).

Red Blood Cell Permeability

RBC water, K⁺ and Na⁺ ion contents were measured following RBC washing in saline buffer containing calcium, 48h after blood collection in EDTA tubes. Figure 2 illustrates data coming from experiments corresponding to two different shipments of either proband A or proband B blood samples collected at different times. Osmotic resistance curves for proband A as for the mother with V222L mutation were slightly shifted to the left compared to control (Figure 2A). However, this shift is in the normal range and there was no change in water content between control and mutant (Figure 2B). The K⁺ and Na⁺ contents of RBC with V222L KCNN4 were not significantly different from control despite a tendency to increased Na⁺ content correlated with decreased K⁺ content in one blood sample (Figures 2C,D). Osmotic resistance curves showed no significant difference between control and H340N mutants (Figure 2E). This is correlated with no significant change in water content (Figure 2F). The K⁺ and Na⁺ contents were also similar between control and H340N mutants (Figures 2G,H). Hence, it was not possible to detect any significant change in water, K⁺, and Na⁺ permeability in RBC carrying KCNN4 mutations V222L or H340N compared to WT. Similar results were observed...
previously in RBC with R352H mutated KCNN4 (Andolfi et al., 2015; Glogowska et al., 2015; Rapetti-Mauss et al., 2015).

In order to unmask a possible difference in the activation of mutated KCNN4 compared to WT, control and mutant RBCs were challenged with vanadate, inhibitor of the Ca\(^{2+}\)-ATPase pump. Blocking Ca\(^{2+}\) pump leads to intracellular Ca\(^{2+}\) increase that in turn activates KCNN4. This activation resulted in K\(^+\) loss that can be blocked by Senicapoc, inhibitor of KCNN4. A significant dehydration correlated with K\(^+\) loss was observed in control RBC 40 min after vanadate addition that was blocked by Senicapoc (Figures 3A,B,E). Bright colors represented Senicapoc effect compared to untreated conditions. Data are given as mean ± SD of triplicate experiments done on either a single blood shipment for proband A [panel (A,B,C)] or two different blood shipments for proband B [panel (D,E,F)]. Fresh RBCs were washed in Ringer without calcium and incubated with 5 μM Fluo-4AM 30min at 37°C. Fluo-4AM-loaded RBCs were then incubated in Ringer solution with 1 mM Ca\(^{2+}\) and 5 mM vanadate. Fluorescence was measured with a FACS as a function of time. Data are given as FITC in arbitrary units, median of a single experiment, 10,000 events. The fluorescence linked to basal intracellular Ca\(^{2+}\) concentration was in arbitrary units, mean ± s.e.m.: 750 ± 9 for proband A, 1,167 ± 6 for proband A’s mother, and 839 ± 6 for control and 528 ± 5 for proband B compared to 533 ± 7 for control.

Intracellular Ca\(^{2+}\) rise following Ca\(^{2+}\) pump inhibition with vanadate depends on the Ca\(^{2+}\) leak allowing progressive Ca\(^{2+}\) uptake. Hence, the absence of differential activation of mutated KCNN4 compared to WT in RBC incubated with vanadate could be explained by variations in intracellular Ca\(^{2+}\) concentration between samples. Using the fluorescent probe Fluo4, intracellular Ca\(^{2+}\) concentration was measured in the

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**FIGURE 3** | RBC ion, water, and Ca\(^{2+}\) contents in response to vanadate. Washed RBCs were submitted to 5 mM Vanadate with or without 4 μM Senicapoc treatment in the presence of 0.5 mM ouabain. Water, K\(^+\) and Na\(^+\) contents were measured at different time points (0’, 16’ and 40’). Water content (A,D), K\(^+\) content (B,E) and Na\(^+\) content (C,F) are represented for control in blue, V222L proband A in red and proband A’s mother in purple; panel (A,B,C) and control in blue, H340N proband B in green; panel (D,E,F). Bright colors represented Senicapoc effect compared to untreated conditions. Data are given as mean ± SD of triplicate experiments done on either a single blood shipment for proband A [panel (A,B,C)] or two different blood shipments for proband B [panel (D,E,F)]. (G,H): Fresh RBCs were washed in Ringer without calcium and incubated with 5 μM Fluo-4AM 30min at 37°C. Fluo-4AM-loaded RBCs were then incubated in Ringer solution with 1 mM Ca\(^{2+}\) and 5 mM vanadate. Fluorescence was measured with a FACS as a function of time. Data are given as FITC in arbitrary units, median of a single experiment, 10,000 events. The fluorescence linked to basal intracellular Ca\(^{2+}\) concentration was in arbitrary units, mean ± s.e.m.: 750 ± 9 for proband A, 1,167 ± 6 for proband A’s mother, and 839 ± 6 for control and 528 ± 5 for proband B compared to 533 ± 7 for control.
different RBC incubated with vanadate (Figures 3G,H). Vanadate increased similarly intracellular calcium concentration in control and patients’ RBC in the different experiments.

**KCNN4 Expression in HEK293T Cells**

To study the effect of KCNN4 mutations on the channel activity, HEK293T cells were transfected with either WT KCNN4 or H340N KCNN4 or V222L KCNN4. It is observed that the maximal current intensity is not changed by the mutations (Figures 4A–D). However, the two mutants V222L and H340N were more sensitive to Ca^{2+} compared to WT with an IC50 = 0.28 and 0.20 µM, respectively (0.99 µM for WT). The expression at the plasma membrane of the different constructs was assessed by western blotting of biotinylated proteins. Immunodetection of E-cadherin was used as a reference for expression of endogenous membrane protein to compare the different conditions. All the constructs were similarly expressed at the plasma membrane (Figures 4E,F), ruling out an effect of the mutations on protein addressing the HEK293T plasma membrane. The mutants were sensitive to Senicapoc, blocker of KCNN4, showing a sensitivity similar to the WT (Figures 4G–K).
The two mutations are located in very different regions of the channel (Figures 5A, B). The multiple sequence analysis also showed drastic different profiles for V222L and H340N mutations in terms of conservation. Indeed, the weblogo established from this alignment (Figure 5C) designates H340 as a highly conserved residue (99%), in contrast the Valine in position 222 is poorly conserved (8%). This last position is actually mainly occupied by Threonine (83%). Leucine residue can be found but extremely rarely in the superfamily of KCNN4 (0.9%).

The 3D structure shows that H340 is located in the binding site with the constitutively bound calmodulin (CaM) C-lobe (Figure 5B). More precisely, H340 is carried by the intracellular HB helix, in the region facing the HA helix and close to the peripheral helices S2–S3 linker. H340 is in contact with residues E321, M324, F325, and H328 of HA and A336 and L343 of HB. V222 is located in the transmembrane part of the channel, in the middle of the interface bridging the S1/S4 peripheral helices with the S5/S6 helices that constitute the pore domain. V222 is in contact with both L154 and L157 situated in the peripheral S4 helix and with the conserved residue V266 from helix S6. It is also close to the interface between S1 and S5 helices involving highly conserved residues, i.e., V41 (87%) and E45 (93%) on one side and residues W221 (99%) and R228 (99%) on the other side.

Preliminary results obtained for the mutated modeled structures relaxed with molecular dynamics simulations did not show major changes in the environment of the residue, most local contacts being preserved. However, this does not preclude long range effects as observed through slight displacements of helices for the two mutants along the molecular dynamic simulations.

DISCUSSION

At the molecular level, it was shown that the KCNN4 mutations V222L and H340N altered the Ca2+ sensitivity of the K+ channel and could be classified as gain-of-function mutations. The IC50 is shifted toward lower intracellular Ca2+ concentrations suggesting that the mutated channels might be activated with a lower Ca2+ increase compared to WT channels. The functional changes observed for the two mutants may originate from two mechanisms: the first one involving V222L could be due to the fine regulation that exists between the peripheral helices and the pore helices during the transport; in contrast, the second one involving H340N may participate in the activation process itself, by changing the interplay between CaM and the channel. In this last case, it is important to note that other mutations (R352, A322, and S314) in CaM C-lobe-binding site have been reported.

Clinically, these two new KCNN4 mutations blurred a bit more the phenotype of KCNN4 gain-of-function mutations. From Table 1, it appears that KCNN4 gain-of-function mutations are associated with more or less compensated hemolytic anemia. For proband A with V222L mutation, the anemia seemed related to dyserythropoiesis and strikingly the mother carrying the same mutation had no history of anemia. For proband B with H340N mutation,
the anemia was transient and again the mother carrying this mutation had never been followed for hematological disorder. The complete response observed under steroids, however, challenges the responsibility of the KCNN4 mutation alone in the pathophysiology of this transient anemia. This questions the role of these two KCNN4 gain-of-function mutations in proband A’s constitutive or proband B’s transient anemia. The hematologic disorder might result from the combination between the presence of KCNN4 gain-of-function mutations and a secondary trigger such as an infection, or a cumulative mutation on another gene involved in erythropoiesis, not included in the 93 "RBC genes" of the t-NGS library, or structural allelic variation not identified in t-NGS. A previous article reported the H340N KCNN4 mutation in a genetic study enrolling 155 patients with suspected hereditary anemia (Andolfo et al., 2021). This KCNN4 mutation was identified in two patients from different kindred and it was associated with either a SPTA1 mutation or two independent mutations on LARS2 and ABCB6. Unfortunately, no hematological or clinical data were given and we could not compare it with the present cases. Considering proband A (V222L mutation), the non-regenerative anemia may be due to either a non-identified intercurrent cause, or a role of KCNN4 during erythropoiesis, as described for PIEZO1-related DHST (Caulier et al., 2020).

At the cell level, neither V222L nor H340N KCNN4 gain-of-function mutations did correlate with altered RBC hydration status: 48 h after withdrawal in EDTA collecting tubes, the osmotic resistance curves were within normal range. When assessed, the MCHC was normal or slightly above the maximal value, and the water, Na⁺, and K⁺ contents were not significantly different from WT. Previous data with other KCNN4 mutations reported by us or others in different publications also failed to correlate KCNN4 mutations with RBC dehydration (Andolfo et al., 2015; Fermo et al., 2015; Rapetti-Mauss et al., 2015; Fermo et al., 2017; Waldstein et al., 2021).

The higher Ca²⁺ sensitivity of H340N and V222L KCNN4 as well as the other characterized mutants S314P and R352H (Rapetti-Mauss et al., 2015; Rivera et al., 2017; Fermo et al., 2020) suggested that these channels would be more often activated in RBC challenged with increasing Ca²⁺ concentration. This higher Ca²⁺ sensitivity of mutated KCNN4 was expected to induce RBC shrinking due to net KCl loss accompanied by water loss that is not compensated in human RBC which does not regulate their volume. It could be argued that to observe KCNN4 activity, Ca²⁺ has to be present in extracellular medium and its concentration should increase in the cell (Dyrda et al., 2010). In ektactometry experiments as in osmotic resistance measurements, there was no Ca²⁺ in extracellular medium and the water uptake or the elongation resulted in no change or at worse a decrease in intracellular Ca²⁺ (due to water uptake). These experimental conditions do not promote KCNN4 activation. We have previously shown that osmotic resistance curves done on RBC carrying R352H KCNN4 mutation treated with the anticoagulant heparin, within hours following withdrawal, were shifted to the left compared to WT, indicating RBC dehydration (Rapetti-Mauss et al., 2015). A similar shift to the left on ektactometry curves done on RBC with R352H or A322V KCNN4 was observed in another laboratory, indicating experimental conditions allowing KCNN4 activation in patient RBC compared to control WT (Mansour-Hendili et al., 2021). Moreover, at the time ektactometry analysis was done on proband B, a shift to the left of the curve indicated dehydration that was corroborated by increased MCHC (Table 1 age 1.3 years). Hence, there might be a situation where the RBC carrying KCNN4 mutations would appear dehydrated. To unmask the consequences of KCNN4 gain-of-function mutations on RBC water homeostasis, Ca²⁺ must be present in extracellular medium and the RBC energy depleted: without ATP, the Ca²⁺ pump activity decreases and intracellular Ca²⁺ slowly increases leading to the activation of KCNN4. Gain-of-function KCNN4 mutations will result in a more rapid activation of the channel compared to WT. Thus, depending on 1) the energy status of RBC following blood withdrawal and 2) the presence of Ca²⁺ in medium, mutated KCNN4 could be activated and patient RBC dehydration could be observed. The fact that in EDTA collecting tubes within 24–48 h of withdrawal, patient RBC was within normal range for water content, suggested KCNN4 activity was kept silent in bloodstream. Alternatively, we can hypothesize that the RBC where KCNN4 was activated were dramatically dehydrated and immediately removed from circulation and could not be observed in blood samples. However, blood smear clearly indicated altered RBC shapes (Figure 1) that were even more visible when patients were splenectomized (Rapetti-Mauss et al., 2015). In addition, the ektactometry indicated in some patients an increased RBC fragility (diminution of the maximum and increase in the minimum elongation index, proband A Figure 1). Nonetheless, these altered shapes or RBC fragility could not be linked to changes in RBC hydration status, or K⁺ and Na⁺ contents. This questions the relevance of connecting the hydration status and RBC fragility in case of KCNN4 gain-of-function mutations.

Moreover, it reinforces the phenotype distinction between KCNN4 gain-of-function mutations and gain-of-function mutations in PIEZO1. In similar experimental condition (EDTA collecting tubes), PIEZO1-mutated RBC is most of the time dehydrated and ektactometry or osmotic resistance curves show a leftward shift of the curves compared to control (Rapetti-Mauss et al., 2017; Picard et al., 2019; More et al., 2020; Andolfo et al., 2021). The RBC dehydration linked to PIEZO1 gain-of-function mutations was explained by a more frequent stimulation of KCNN4 (Albuissou et al., 2013), leading to a new RBC volume homeostasis. The role of PIEZO1 in RBC volume homeostasis was confirmed by the observation that mice or zebrafish RBC knockdown for piezo1 was hyperhydrated compared to WT RBC (Faucherre et al., 2014; Cahalan et al., 2015). Moreover, a patient with combined PIEZO1 mutations correlated with decreased expression of the protein exhibited overhydrated RBC (Andolfo et al., 2015).

The present data combined with the previous publications reinforce the difficulty to link hemolytic anemia to KCNN4 gain-of-function mutations. The activity of the mutated channel appeared to be most of the time under strict control, kept silent in RBC with one notable exception, the V282M mutation that yields a constitutive K⁺ leaky channel that is clearly associated with RBC dehydration (Rivera et al., 2017). Since 2015, and taking into account the new mutations reported
here, 11 different KCNN4 mutations have been identified in independent kindreds (P204R, A279T, S314P, V222L, V282/M or E, H340N, R352H, and delV369-L373) (Andolfo et al., 2015; Glogowska et al., 2015; Rapetti-Mauss et al., 2015; Ferro et al., 2017; Utsugisawa et al., 2017; Picard et al., 2019; Ferro et al., 2020; Andolfo et al., 2021; Mansour-Hendili et al., 2021). According to the different patients studied so far, using the RBC hydration status to diagnose a DHSt associated to KCNN4 mutations is perhaps misleading. The dehydration seems to be a signature of PIEZO1 gain-of-function mutations and exceptionally of KCNN4 mutation, misleading. The dehydration seems to be a signature of PIEZO1 gain-of-function mutations and exceptionally of KCNN4 mutation, when the mutation converts the channel into a constitutive K+ leak (the V282M mutation is the only one documented so far). Nonetheless, the presence of a gain-of-function mutated Gardos channel in RBC can dramatically impair RBC viability by a mechanism that remains to be identified. This fragility might involve other KCNN4 functions than permeability or require additional events. The susceptibility for a given KCNN4 gain-of-function mutation to lead to hemolysis appears more dramatic in early life, which is a classical feature of constitutional RBC membrane hemolytic anemia. More experimental data will be needed to better understand the physiological role of KCNN4 and how the mutations alter its function in RBC.

**DATA AVAILABILITY STATEMENT**

The data presented in the study are deposited in the SRA repository, accession number PRJNA853732.

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**ETICS STATEMENT**

The studies involving human participants were reviewed and approved by Comité de protection des personnes Île de France. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

**AUTHOR CONTRIBUTIONS**

BA, LDN, MM, RR-M, OF, LDC, and HG did the experiments on blood samples. SJ and CE performed the bioinformatic analysis of the channel. AL supervised the calcium fluorometry. AB, CT, JD, and AP were the clinicians who followed the patients. All the authors analyzed the data and reviewed the manuscript.

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