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Zinc-dependent multimerization of mutant calreticulin is required for MPL binding and MPN pathogenesis

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Key Points

- Zinc is required for multimerization of mutant calreticulin, which is a requisite event for MPL binding and activation.
- Zinc chelators exhibit selective toxicity against CALR-mutant MPN cell lines and primary MPN cells.

Calreticulin (CALR) is mutated in the majority of JAK2/MPL-unmutated myeloproliferative neoplasms (MPNs). Mutant CALR (CALR^{del52}) exerts its effect by binding to the thrombopoietin receptor MPL to cause constitutive activation of JAK-STAT signaling. In this study, we performed an extensive mutagenesis screen of the CALR globular N-domain and revealed 2 motifs critical for CALR^{del52} oncogenic activity; (1) the glycan-binding lectin motif and (2) the zinc-binding domain. Further analysis demonstrated that the zinc-binding domain was essential for formation of CALR^{del52} multimers, which was a co-requisite for MPL binding. CALR^{del52} variants incapable of binding zinc were unable to homomultimerize, form CALR^{del52}-MPL heteromeric complexes, or stimulate JAK-STAT signaling. Finally, treatment with zinc chelation disrupted CALR^{del52}-MPL complexes in hematopoietic cells in conjunction with preferential eradication of cells expressing CALR^{del52} relative to cells expressing other MPN oncogenes. In addition, zinc chelators exhibited a therapeutic effect in preferentially impairing growth of CALR^{del52}-mutant erythroblasts relative to unmutated erythroblasts in primary cultures of MPN patients. Together, our data implicate zinc as an essential cofactor for CALR^{del52} oncogenic activity by enabling CALR^{del52} multimerization and interaction with MPL, and suggests that perturbation of intracellular zinc levels may represent a new approach to abrogate the oncogenic activity of CALR^{del52} in the treatment of MPNs.

Introduction

Mutations in *CALR*, which encodes the endoplasmic reticulum-resident chaperone protein calreticulin (CALR) are present in the majority of *JAK2/MPL*-unmutated myeloproliferative neoplasms (MPNs).^{1,2} CALR is a calcium-binding chaperone that regulates cellular proteostasis in the endoplasmic reticulum.³ Wild-type CALR is composed of 3 domains: (1) a globular *N*-domain that contains lectin-, polypeptide-, and zinc-binding motifs that enable CALR chaperone activity⁴⁻⁶; (2) a P-domain that forms a hairpin structure that binds to cochaperones^{7,8}; and (3) an acidic C-terminal tail involved in calcium buffering.⁹ MPN-associated *CALR* mutations are insertion-deletion mutations that result in a +1-bp frameshift that generates a neomorphic C-terminal tail encoded in an alternative reading frame, with the most common mutation being a 52-bp deletion.^{1,2} The resulting mutant CALR protein (CALR^{del52}) contains

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Data and reagents will be made available to other investigators upon request to the corresponding author at e.chen@westminster.ac.uk.

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a mutant-specific C terminus in which most of the negatively charged amino acids of the wild-type protein is replaced with positively charged amino acids, such as lysine and arginine.

Multiple studies have now provided direct links between CALR^{del52} and thrombopoietin receptor (MPL) signaling. Expression of CALR^{del52} confers cytokine-independent growth in conjunction with increased JAK-STAT signaling in Ba/F3 and UT7 cell lines coexpressing MPL, but not erythropoietin (EPO) receptor or granulocyte-macrophage colony-stimulating factor receptor.¹⁰⁻¹² Consistent with an essentiality of MPL in mediating CALR^{del52} -associated disease, CALR^{del52} was unable to induce MPN-like disease in mice lacking *Mpl.*¹³ Mechanistic studies revealed that CALR^{del52} physically associates with the extracellular domain of MPL¹⁰⁻¹² through its lectin motif^{10,14} to activate JAK-STAT signaling and engender cytokine-independent proliferation. Moreover, CALR^{del52} has also been shown to homomultimerize, and that this homomultimerization is a prerequisite for MPL binding and activation.¹⁵

In this report, we undertook a comprehensive mutagenesis screen and uncovered a new role for 3 zinc-binding histidine residues in the CALR^{del52} *N*-domain essential for CALR^{del52} multimerization and MPL binding and activation. Furthermore, we found that the CALR^{del52}-MPL signaling complex is dependent on intracellular zinc. We show that zinc chelators were able to disrupt CALR^{del52} -MPL signaling complexes and attenuate JAK-STAT signaling specifically in CALR-mutant cells, and exhibited selective toxicity against CALR^{del52}-positive burst-forming unit erythroid (BFU-E) colonies from MPN patients. Our data reveal several additional insights into the molecular mechanism of action of CALR^{del52} by identifying a central role for zinc in regulating CALR^{del52} multimerization and its impact on MPL signaling.

Materials and methods

Cell culture reagents and antibodies

The 293T cells were derived from the American Type Culture Collection. MPL-expressing Ba/F3 (Ba/F3-MPL) were generated by retroviral transduction with retroviruses expressing human MPL complementary DNA. The 293T cells were maintained in Dulbecco's modified Eagle medium (DMEM; Lonza), supplemented with 10% fetal calf serum (Sigma) and 1% penicillin-streptomycinglutamine mix (Sigma), and cultured in 37°C with 5% CO₂. Ba/F3-MPL cells were maintained in RPMI (Lonza) supplemented with 10% fetal calf serum, 1% penicillin-streptomycin-glutamine mix, and 50 ng/mL recombinant human thrombopoietin (PeproTech) in a vented flask, and cultured in 37°C with 5% CO2. For N,N,N',N'tetrakis(2-pyridinylmethyl)-1,2-ethanediamine (TPEN) treatment, TPEN (Sigma) was resuspended in 99% ethanol at a concentration of 20 µM and added to media at indicated concentrations for 1 to 4 hours. For clioquinol treatment, clioquinol (Sigma) was resuspended in dimethyl sulfoxide at a concentration of 40 µM and added to media at indicated concentrations for 4 hours to assess effect intracellular signaling and 48 hours to assess cell viability. For immunoblotting, primary antibodies were obtained from the following sources: mouse anti-FLAG-M2 (Sigma), rabbit anti-FLAG rabbit (Sigma), rabbit-anti-V5 (Cell Signaling), rabbit anti-CALR (clone D3E6) (Cell Signaling), rabbit anti-pSTAT5-Y694 (clone D47E7) (Cell Signaling), rabbit anti-pSTAT3-Y705 (clone D3A7) (Cell Signaling), rabbit anti-total STAT5 (clone D2O6Y) (Cell signaling), rabbit anti-STAT3 (clone124H6) (Cell Signaling), mouse anti-ACTIN (clone AC35) (Sigma), and mouse anti-MPL (clone 1.78.1) (BD Bioscience).

Alanine mutagenesis screen

Mutagenesis of iV2-LeGO-iV2-CALR^{del52} plasmid DNA was performed using Q5 Hot Start High-Fidelity Master mix (New England Biolabs) according to the manufacturer's protocols. The following thermocycling setup was used: initial denaturation 98°C for 30 seconds, 25 cycles at 98°C for 10 seconds, 50°C to 72°C for 20 seconds, and 72°C for 4 minutes, followed by a final extension at 72°C for 10 minutes. Kinase, ligase, and DpnI treatment was immediately performed following the linear amplification for 1 hour at 37°C. The kinase, ligase, and DpnI-treated reaction was used to transform competent Escherichia coli. DNA was isolated from individual colonies, and presence of alanine mutant was confirmed by Sanger sequencing. To perform functional assays for each CALR alanine variant, lentiviral supernatants were prepared for each alanine variant by cotransfection of the mutagenized LeGO-iV2-CALR variant with packaging plasmids psPAX2 and pVSV.G. After 24 hours of transfection, CALRexpressing viral supernatants were collected, filtered, and used to infect Ba/F3-MPL cells by spin infection at 2000 rpm at 37°C for 2 hours. Cells were allowed to recover by incubation with RPMI supplemented with 50 ng/mL recombinant human thrombopoietin (PeproTech) for 2 days, before cytokine withdrawal. For cytokine withdrawal, exponentially growing Ba/F3 MPL cells transduced with lentiviruses expressing various CALR variants were washed 3 times with phosphate-buffered saline (PBS) and seeded in triplicate at an initial density of 2×10^5 cells/mL in RPMI-1640 medium (Lonza) supplemented with 10% fetal bovine serum (Sigma) and 1% penicillin/streptomycin/L-glutamine (Sigma) in the absence of cytokine. Living cells were counted using hemocytometers by trypan blue exclusion assay. 3-(4.5-dimethylthiazol-2-vl)-2.5-dimethyltetrazolium bromide assays or by cell counting on a flow cytometer. The ability of each alanine variant to confer cytokine-independent growth was depicted relative to that of an unmutated LeGO-iV2-CALR^{del52} control.

FLAG pulldown assays

To assess protein complex formation of heterologously expressed proteins, coimmunoprecipitations were performed in 293T cells. Cells were seeded at a density of 2×10^5 cells/mL in a 6-well plate. For assessing binary CALR^{del52}-MPL complex formation, DNAcationic lipid mixtures were created by combining 1.25 µg of pCMV-SPORT6-CALR variant-expressing plasmid, 1.25 µg of pLeGO-iT2-hMPL, and 7.5 µL TransIT-LT1 transfection reagent (Mirus) in 250 µL of media-free DMEM. For assessing ternary CALR^{del52}-MPL complexes with 2 different tagged versions of CALR^{del52}, 0.625 μg of each pCMV-SPORT6-CALR variantexpressing plasmid was used and mixed with 1.25 µg of pLeGOiT2-hMPL and 7.5 µL TransIT-LT1 transfection reagent in 250 µL media-free DMEM, as discussed previously. Mixtures were allowed to sit at room temperature for 15 minutes, and then added dropwise to 293T cells. After 24 hours, transfected 293T cells were harvested, washed with PBS twice, and then cells were collected and resuspended in 450 μ L NP40 lysis buffer and incubated on ice for 30 minutes. Lysates were cleared by centrifugation at 17000g for 10 minutes at 4°C, and 10% (usually \sim 50 μ L) was set aside as the total lysate "input" fraction and the remainder used for pulldown assays. Protein G bead-antibody mixtures were generated by initially washing 50 μ L of magnetic Protein G Surebeads (BioRad Laboratories) per sample with PBS with 0.1% Tween-20 [PBST]) 3 times using a magnet. One microliter of FLAG M2 antibody (Sigma) was diluted with PBST and added to the beads and incubated for 15 minutes with inversion at room temperature. FLAG M2-Protein G beads were washed with PBST and incubated with lysate supernatants for 1 hour at room temperature and mixed by inversion. Lysate-bead mixtures were washed 3 times by magnetic sorting, and beads were resuspended in 100 μ L of 1X SDS Laemmli buffer and eluted by incubating the beads for 10 minutes at 70°C. Both pulldown fractions and lysates were analyzed by Western immunoblotting, as described in supplemental Information.

Intracellular phosphoprotein flow cytometry analysis

Ba/F3-MPL cells expressing CALR variants were cultured in RPMI supplemented with 10% fetal bovine serum, 5% penicillin/ streptomycin, and 5% L-Glutamine for 24 hours in the absence of cytokines. Harvested cells were transferred into flow tubes and washed with PBS + 1% bovine serum albumin (BSA) twice to remove residual RPMI media. These cells were fixed and permeabilized using Fix&Perm Kit (Nordic MUbio). Staining was carried out in PBS 1% BSA for 1 hour at room temperature with PE Mouse anti-Stat3 (pY705) and Alexa Fluor 647 anti-Stat5 (pY694) (BD Biosciences). Cells were washed with PBS with 1% BSA and samples were analyzed using a CytoFLEX Flow Cytometer (Beckman Coulter Life Sciences). Data analysis was performed with CytExpert V2.2.

Confocal microscopy and FRET analysis

The 293T cells were seeded into 35mm, high, μ -Dish with an ibidi polymer coverslip bottom and were incubated overnight at 37°C. Transient co-transfection of pSport6-Cherry-CALR variant-FLAG with MSCV-GFP-MPL in a 1:1 ratio was performed using TransIT-LT1 (Mirus). Following 24 hours post-transfection, cells were gently washed and resuspended with phenol red-free DMEM media, and cells were imaged using Zeiss LSM 880 inverted confocal microscopy. Fluorescence resonance energy transfer (FRET) was measured by exciting sample at donor excitation of 488 nm and measuring the acceptor excitation absorbance at 610 nm. FRET intensity was calculated by using sensitized emission.

TPEN treatment of clonal erythroblasts from MPN patients

MPN patients were diagnosed according to the British Committee for Standards in Hematology guidelines.¹⁶ UK patient samples were obtained under local ethics approval (Cambridge and Eastern Region Ethics Committee) or as part of the UK Medical Research Council PT-1 trial.¹⁷ Written informed consent was obtained from all patients in accordance with the Declaration of Helsinki. For local Cambridge patients, samples were taken at the time of the initial patient visit on referral to the specialist MPN clinic at the Addenbrooke's Hospital in Cambridge, United Kingdom. Venous blood samples (20 mL) were collected from each patient and peripheral blood mononuclear cells (PBMCs) were isolated using Lymphoprep (Axis Shield PLC) according to the manufacturer's protocols and plated in complete MethoCult medium (H4435; STEMCELL Technologies) at a density of 3×10^5 cells/mL and incubated at 37°C and 5% CO₂ for 14 days. When indicated, MethoCult was supplemented with 5 μ M TPEN (Sigma). After 14 days, colonies were plucked into 20 μ L PBS. For each colony, 5 μ L was denatured at 95°C for 15 minutes to release cellular DNA and subjected to polymerase chain reaction using primers to amplify exon 9 of the *CALR* gene (forward primer: 5'-GCA GCA GAG AAA CAA ATG AAG G-3'; reverse primer: 5'-AGA GTG GAG GAG GGG AAC AA-3'). Polymerase chain reaction conditions were an initial preheating at 94°C for 10 minutes was followed by denaturation at 94°C for 30 seconds, annealing at 64°C for 30 seconds, and extension at 72°C for 7 minutes. Amplicons were submitted for Sanger sequencing to identify colony genotypes.

Results

Alanine mutagenesis screen of CALR^{del52} *N*-domain demonstrates residues associated with lectin and zinc-binding activity are essential for oncogenic activity

We and others previously reported the lectin motif within the CALR^{del52} N-domain was essential for MPL binding.^{10,14} To resolve other regions within the N-domain potentially necessary for oncogenic activity, we undertook an alanine mutagenesis screen of the entire *N*-domain by screening for the ability of single-residue loss-of-function variants to engender cytokine-independence in MPL-expressing Ba/F3 (Ba/F3-MPL) cells. We observed that the majority of the 177 single-residue alanine mutants did not affect CALR^{del52} activity, with only 9 residues abrogating cytokine independence by a factor greater than 1 standard deviation from the mean when mutated (Figure 1A). Specifically, CALR^{del52} harboring C105A, K111A, G133A, or D135A mutations exhibited total abrogation of cytokine independence (Figure 1B, left), decreased STAT3/5 phosphorylation as assessed by intracellular flow cytometry (Figure 1C) and immunoblotting (Figure 1D), and an inability to bind MPL in FLAG-pulldown assays (Figure 1E). These residues have previously been implicated in the lectin activity of wild-type CALR,18 and D135 specifically has previously been shown by us and others to be required for CALR^{del52} oncogenic activity.^{10,14} In addition, CALR^{del52} harboring Y109A and Y128A mutations only exhibited partial loss of cytokine independence (Figure 1B, right) and exhibited no effect on STAT3/5 phosphorylation or MPL binding (Figure 1C-E), despite these residues also having been implicated in wild-type CALR lectin activity. Of note, other residues thought to be essential for CALR lectin motif such as M131 and I147 did not exhibit any effect in our screen (Figure 1A). Overall, these data accord with previous findings that the lectin motif of CALR^{del52} is essential for oncogenic activity, but suggest that different residues within the lectin motif of CALR^{del52} may have differential effects in facilitating its oncogenic activity.

Next, we examined the other residues identified in our screen that have not previously been associated with glycan binding. We observed that CALR^{del52} harboring H99A, H145A, and H170A mutations (1His) exhibited partial abrogation of cytokine independence (Figure 1A and Figure 2A, left). Because these residues have been reported to need to function in a concerted manner to coordinate zinc,¹⁹ we tested whether combined loss of multiple residues would have more deleterious consequences on CALR^{del52} activity. Indeed, we observed that loss of any 2 or all 3 histidine residues in combination (2His and 3His, respectively) led to

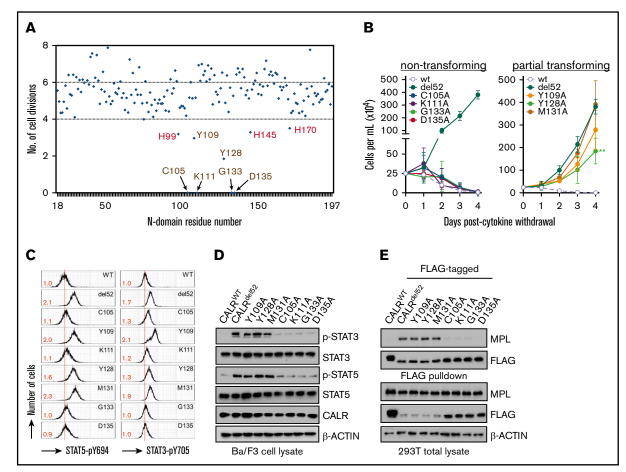


Figure 1. The lectin-dependent function of CALR^{del52} **is required for cytokine-independent growth.** (A) Alanine mutagenesis screen to assess ability of 177 CALR^{del52} single-residue mutants to confer cytokine-independent growth of Ba/F3-MPL cells, as depicted on y-axis by number of cell divisions following 3 days. The mutagenized residue is represented on the x-axis. The 9 residues that are >1 standard deviation away from mean are indicated with arrows. Putative lectin residues are denoted in brown; putative zinc-binding residues are denoted in red. (B) Growth curves of Ba/F3-MPL cells expressing wild-type CALR, CALR^{del52}, or lectin motif variants demonstrate total (left) or partial (right) impairment of cytokine-independent growth. Testing for statistical significance was performed using a Student *t* test (**P* < .05; ***P* < .01; ****P* < .001). (C) Intracellular phosphorylation flow cytometry and (D) Western immunoblotting demonstrate diminished Stat3 and Stat5 phosphorylation in Ba/F3-MPL cells expressing nontransforming CALR^{del52} lectin variants. Numbers in red indicate ratio of mean fluorescence intensity for each sample relative to isotype control. The data are representative of 2 independent experiments. (E) Immunoblotting of FLAG-immunoprecipitated proteins from 293T cells cotransfected with FLAG-tagged CALR^{del52} lectin variants and MPL-expressing vector demonstrates impaired MPL binding capacity by nontransforming lectin variants.

significant impairment of cytokine-independent growth (Figure 2A, middle and right), in conjunction with decreased pSTAT3/5 phosphorylation (Figure 2B-C).

To ensure that these results are specific to these individual zincbinding histidine residues and not from nonspecific effect on CALR^{del52} stability, we also tested the effects of CALR^{del52} in which H42, which is predicted to lie on the opposite face of CALR^{del52} and is not thought to function in concert with H99, H145, or H170 to bind zinc,¹⁸ is mutated. We observed that CALR^{del52} variants harboring H42A mutations or of H42A in combination with H99, H145, or H170 were unaffected in their ability to confer cytokineindependent growth (supplemental Figure 1A) and retained the capacity to promote STAT3/5 phosphorylation (supplemental Figure 1B-C).

Finally, we also tested whether histidine-deficient CALR^{del52} variants could physically associate with MPL by co-immunoprecipitation assays. We observed that 1His-CALR^{del52} variants retained the

ability to bind to MPL in FLAG pulldowns, but not 2His or 3His variants (Figure 2D). We also tested the ability of histidinedeficient CALR^{del52} variants to colocalize intracellularly with MPL using FRET. To achieve this, histidine-deficient CALR^{del52} variants were generated fused to an mCherry fluorescent marker and introduced to 293T cells in conjunction with an MPL-GFP fusion protein; colocalization was measured by quantification of energy transfer of the mCherry donor to the GFP acceptor. Significant FRET was observed for mCherry-tagged CALR^{del52} and MPL-GFP indicative of intracellular colocalization, whereas minimal FRET was observed for mCherry-tagged wild-type CALR and MPL-GFP (supplemental Figure 2). Analysis of 100 cells for each histidine-deficient CALR^{del52} variants indicated that MPL colocalized strongly with 1His-CALR^{del52} variants, but not with 2His and 3His variants (Figure 2E).

Cumulatively, functional analysis of single-residue CALR^{del52} *N*-domain variants were able to identify 2 classes of residues that

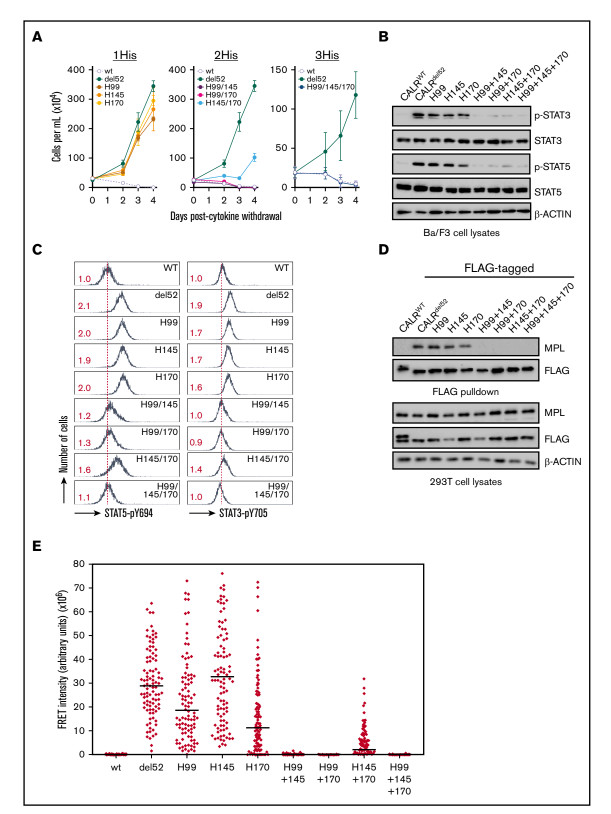


Figure 2. A triad of zinc-binding histidine residues in CALR^{del52} are required for cytokine-independent growth. (A) Growth curves in Ba/F3-MPL cells expressing histidine-deficient CALR^{del52} variants harboring loss of either 1 histidine (1His), 2 histidines (2His), or 3 histidines (3His). Immunoblotting (B) and intracellular phospho-flow cytometry (C) demonstrate diminished Stat3 and Stat5 phosphorylation in of Ba/F3-MPL cells expressing 2His- and 3His-CALR^{del52} variants. Numbers in red indicate ratio of mean fluorescence intensity for each sample relative to isotype control. The data are representative of 2 independent experiments. (D) Immunoblotting of FLAG-immunoprecipitated proteins from 293T cells cotransfected with histidine-deficient FLAG- CALR^{del52} variants and MPL-expressing vectors demonstrates abolished MPL bind-ing capacity by 2His- and 3His-CALR^{del52} variants. (E) Single-cell data for FRET intensity. FRET fluorescence units depict energy transfer between mCherry-tagged histidine-deficient CALR^{del52} protein and MPL-GFP in arbitrary fluorescence units for a single cell. For each experimental condition, 100 cells were analyzed.

are important for mediating the ability of CALR^{del52} to bind and activate MPL and confer cytokine independence: (1) residues associated with lectin motif and (2) histidine residues putatively associated with zinc binding.

Histidine-mutated CALR^{del52} is incapable of homomultimerization

A recent study detected CALR^{del52} homomultimerization by pulldown of distinct CALR^{del52} molecules tagged with 2 different epitopes.¹⁵ We used the same strategy to test the multimerization capacity of lectin-deficient or histidine-mutated CALR^{del52}. We confirmed these findings by assessing the ability of FLAG-tagged CALR^{del52} (bait) to complex and co-immunoprecipitate with a V5tagged CALR^{del52} species (prey) in 293T cells (Figure 3A), and demonstrated that CALR^{del52} is capable of interacting with CALR^{del52} but exhibited impaired ability to bind to wild-type CALR (Figure 3B). Using this experimental setup, we next tested the multimerization capacity of lectin-deficient or histidine-mutated CALR^{del52}. We observed that the lectin-deficient CALR^{del52} and all 3 1His-CALR^{del52} variants retained multimerization capacity equivalent to that of CALR^{del52} (Figure 3B-C). In contrast, we observed that the 2His- and 3His-CALR^{del52} were impaired in their capacity to multimerize with CALR^{del52} (Figure 3C). To ensure that these activities were not an artifact of assessing binding with unmutated CALR^{del52}, we also tested the ability of lectin-deficient or histidine-mutated CALR^{del52} to multimerize with another lectindeficient or histidine-mutated CALR^{del52}, respectively, and form homomultimers. Consistent with our earlier data, we observed that the lectin-deficient CALR^{del52}-D135A retained the ability to homomultimerize with another CALR^{del52}-D135A, but the histidine-mutated 3His-CALR^{del52} was not able to homomultimerize with another 3His-CALR^{del52} (Figure 3D). Our data therefore accord with previous reports that CALR^{del52} forms homomultimers, but our data show that homomultimerization is necessary but not sufficient for MPL binding.

Zinc is essential for CALR^{del52}-MPL complex formation and downstream signaling

H99, H145, and H170 have previously been implicated in zincbinding capacity of wild-type CALR.¹⁹ Therefore, to confirm that loss of zinc coordination underpinned the loss of oncogenic capacity of 2His- and 3His-CALR^{del52}, we performed affinity chromatography using a zinc resin and found that CALR^{del52} and 1His-CALR^{del52} could readily be retained in the zinc-bound fraction, but that wild-type CALR, 2His-, and 3His-CALR^{del52} exhibited diminished zinc binding (Figure 3E). That wild-type CALR did not bind to the zinc resin is somewhat surprising because wild-type CALR has previously been shown to bind zinc.¹⁰ This may suggest that the assay used in our study was not sufficiently sensitive to detect the wild-type CALR-zinc interaction, and that there may be differences in the avidity of wild-type and mutant CALR monomers to zinc and/or that CALR^{del52} (but not wild-type CALR) forms multimers or aggregates when bound to zinc, resulting in greater recovery upon elution. Cumulatively, these data are consistent with the finding that 2His- and 3His-CALR^{del52} exhibit loss of zinc binding.

These data therefore suggested that binding to zinc could be required for CALR^{del52} multimerization and MPL binding. To test this, FLAG- and V5-tagged CALR^{del52} and MPL were expressed in

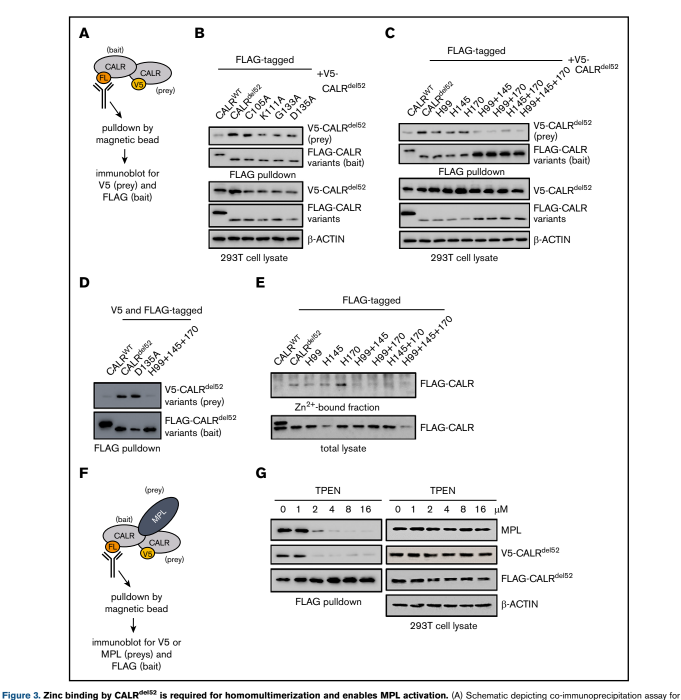
293T cells, treated with agents that modulate zinc levels and FLAG pulldown was performed to assess the extent of CALR^{del52} multimerization and MPL binding in heteromeric complexes (Figure 3F). To modulate zinc levels, TPEN was used as a membrane-permeable ion chelator with a high affinity for zinc.²⁰ Treatment with TPEN led to decreased CALR^{del52} multimerization and impaired MPL binding (Figure 3G), suggesting that zinc is required for CALR^{del52} to multimerize and interact with MPL.

These data would predict that zinc chelation could abolish cytokine independence and JAK-STAT signaling specifically in CALR^{del52} -expressing hematopoietic cells but potentially not in cells expressing other MPN oncogenes. We therefore tested the effect of TPEN on cytokine-independent growth of CALR^{del52}-expressing Ba/F3-MPL cells (Ba/F3-MPL-CALR^{del52}), and observed that TPEN treatment led to increased cytotoxicity (Figure 4A) and attenuated STAT3/5 phosphorylation (Figure 4B; supplemental Figure 3), which were correlated with decreased interaction between CALR^{del52} and MPL (Figure 4C). Moreover, the effect on JAK-STAT signaling was specific to CALR^{del52} because Ba/F3 cells transformed with the alternate MPN oncogene MPL^{W515L} (Ba/F3-MPL^{W515L}) was less sensitive to TPEN-induced cytotoxicity and reduction of STAT3/5 phosphorylation (Figure 4A-B; supplemental Figure 3), despite equivalent reduction of intracellular zinc (supplemental Figure 4A-C). Similar results were obtained using a cytokinedependent megakaryocytic cell line UT7 coexpressing MPL. TPEN induced CALR^{del52}-specific cytotoxicity in UT7 cells (supplemental Figure 5A), disassociation of MPL-CALR^{del52} complexes (supplemental Figure 5B), and selective abrogation of JAK-STAT signaling in CALR^{del52}-expressing cells (supplemental Figure 5C).

Next, to ensure this effect was generalizable to other zinc chelators, we also tested whether clioquinol, which has also been reported to have zinc-chelating activity,²¹ had a similar effect. As with TPEN, treatment of cells with clioquinol also led to preferential CALR^{del52} cytotoxicity in Ba/F3-MPL cells (Figure 4D) and abrogation of JAK-STAT signaling (Figure 4E). Moreover, clioquinol could also promote disassociation of MPL-CALR^{del52} complexes in 293T cells (Figure 4F) These data suggest that sensitivity to zinc homeostasis is a feature of CALR^{del52}-induced MPN.

TPEN preferentially eradicates CALR-mutant primary MPN cells

Finally, we tested the effect of zinc chelation on CALR-mutant primary MPN cells. We initially performed tests to identify TPEN concentrations which would affect growth of wild-type BFU-Es grown from PBMCs in semisolid medium, and observed no impaired growth of wild-type colonies at TPEN dosages lower than at 3 µM (Figure 5A). Next, PBMCs from 5 CALR^{del52}-mutated myelofibrosis (MF) patients (with allele burdens ranging from 20% to 73%) were grown in semisolid medium supplemented with 2.5 µM TPEN. After 14 days, BFU-Es were plucked and genotyped for the presence of CALR mutation by Sanger sequencing. In 4 of 5 MF patients examined, TPEN treatment led to decreased growth of CALR^{del52} -positive BFU-E colonies compared with autologous wild-type colonies (Figure 5B), with the fraction of mutant colonies in TPENtreated cultures reduced to 74% compared with untreated cultures (P = .053) (Figure 5C). The effect of TPEN appeared to be specific to CALR-mutant cells because no selective effect was observed on



Produce 3. Zinc binding by CALR and its required for noncommutimerization and enables MPL activation. (A) Schematic depicting co-immunoprecipitation assay for detecting FLAG-tagged and V5-tagged CALR^{del52} multimerization and enables MPL activation. (A) Schematic depicting co-immunoprecipitation assay for detecting FLAG-tagged and V5-tagged CALR^{del52} multimerization and enables MPL activation. (A) Schematic depicting co-immunoprecipitation assay for detecting FLAG-tagged and V5-tagged CALR^{del52} multimerization and enables MPL activation. (A) Schematic depicting co-immunoprecipitation assay for detecting to V5-CALR^{del52}. (C) FLAG-pulldown assays demonstrating CALR^{del52} lectin variants retain ability to bind to V5-CALR^{del52}. (D) FLAG-pulldown assays of cells transfected with FLAG- and V5-tagged CALR^{WT} (lane 1), CALR^{del52} (lane 2), CALR^{del52} D135A lectin variant (lane 3), or 3His-CALR^{del52} variant (lane 4) demonstrating that CALR^{del52} and CALR^{del52} D135A are capable of homomultimerization, but not CALR^{WT} and 3His-CALR^{del52}. (E) Affinity chromatography demonstrating 2His- and 3His-CALR^{del52} bind less efficiently to zinc resin compared with 1His-CALR^{del52}. (F) Schematic depicting co-immunoprecipitation assay for detecting heteromeric complexes comprising FLAG-tagged and V5-tagged CALR^{del52} proteins and MPL. (G) FLAG-pulldown assays demonstrating zinc chelator TPEN treatment disrupts CALR^{del52} multimerization and MPL binding in 293T cells.

JAK2V617F-mutant BFU-Es in response to TPEN in 4 JAK2-mutant MF patients examined (Figure 5D-E), and no CALR^{del52} mutantspecific eradication was observed following treatment with an established MPN treatment, hydroxyurea (Figure 5F-G). Cumulatively, these data provide a proof of principle that zinc chelation selectively affects the growth of CALR^{del52}-positive patient-derived BFU-Es and has the potential to be harnessed as a therapeutic modality in MPN patients.

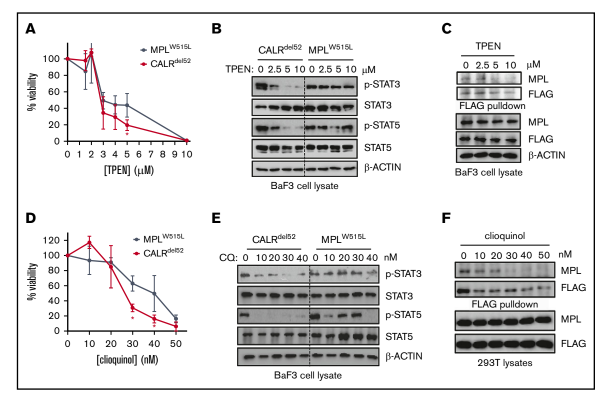


Figure 4. Zinc chelation abrogates CALR^{del52} **-induced JAK-STAT signaling in hematopoietic cell lines.** (A) Viability of BaF3-MPL cells expressing CALR^{del52} or MPL^{W515L} following treatment with TPEN for 48 hours. Each point represents the mean of 3 independent cultures. Testing for statistical significance was performed using a Student *t* test (**P* < .05). (B) Immunoblotting demonstrates decreased Stat3 and Stat5 phosphorylation status following TPEN treatment of 4 hours in CALR^{del52}-expressing Ba/F3-MPL cells but not in Ba/F3-MPL cells. (C) FLAG-pulldown assays demonstrating zinc chelator TPEN treatment disrupts CALR^{del52}-MPL binding in CALR^{del52} -expressing Ba/F3-MPL cells. (D) Viability of BaF3-MPL cells expressing CALR^{del52} or MPL^{W515L} following treatment with clioquinol (CQ) for 48 hours. Cell viability was quantified by 3-(4,5-dimethylthiazol-2-yl)-2,5-dimethyltetrazolium bromide assays. Each point represents the mean of 3 independent cultures. The data are representative of at least 2 independent experiments. Testing for statistical significance was performed using Student *t* test (**P* < .05). (E) Immunoblotting demonstrates decreased Stat3 and Stat5 phosphorylation status following CQ treatment of 4 hours in CALR^{del52}-expressing Ba/F3-MPL cells but not in Ba/F3-MPL cells. (F) Immunoblotting of FLAG-immunoprecipitated proteins in 293T cells expressing CALR^{del52} and MPL demonstrates disruption of CALR^{del52}-MPL binding following CQ treatment of 4 hours.

Discussion

Multiple studies have now provided evidence that *CALR* mutations coopt the MPL-JAK2 signaling axis to engender an MPN phenotype and that this is achieved by direct interaction between CALR^{del52} and the MPL extracellular domain.¹⁰⁻¹² In this report, we attempt to provide further molecular insights into how the CALR^{del52}-MPL binding event is governed. We identified a triad of zinc-binding histidine residues within the CALR^{del52} *N*-domain, which is required for both CALR^{del52} homomultimerization and MPL binding, and for engendering the MPN phenotype. To our knowledge, our report is the first demonstrating of the importance of zinc as an essential cofactor in mediating MPN oncogenic signaling.

The precise mechanism by which zinc enables CALR^{del52} multimerization and MPL activation remains unclear. It was somewhat surprising that 2His- and 3His-CALR^{del52} molecules were incapable of binding to MPL despite the presence of an intact lectin motif. We can envision 2 potential explanations for this finding. One possibility is that zinc binding is required to ensure proper conformation of the lectin motif, and that the absence of zinc renders the lectin motif nonfunctional. Our data point toward a second possibility, whereby CALR^{del52} cannot bind to MPL as a monomer but rather needs to engage MPL as a multimer, and that zinc is necessary to mediate this multimerization. Under this scenario, binding by CALR^{del52} to MPL could require both an intact lectin motif and the capacity for zincdependent homomultimerization (Figure 5H). Indeed, there is precedent for a role for ligand multimerization in cytokine signaling. For example, EPO has been found to form high-molecular-weight species when purified²² and EPO dimers have been shown to exhibit more than 26-fold higher biological activity than its monomeric form.²³ Finally, it is possible that these 2 possibilities are not mutually exclusive because mutant CALR multimerization could affect the activity of the lectin motif through an unknown mechanism to enhance its ability to bind MPL. Atomic level resolution of the structure CALR^{del52}-MPL complex will likely be required to resolve these outstanding issues.

Finally, our findings have potential therapeutic implications. Our data show that chemical chelation of intracellular zinc by 2 different heavy metal chelators was able to promote disassembly of the CALR^{del52}-MPL complex and cause attenuation of JAK-STAT signaling in *CALR*-mutant cells. Moreover, zinc chelation led to growth inhibition of CALR^{del52}-positive erythroblasts from primary MF samples. These data are, to our knowledge, the first to implicate heavy metal homeostasis in MPN pathogenesis and have the potential to support the development or repurposing of zinc or

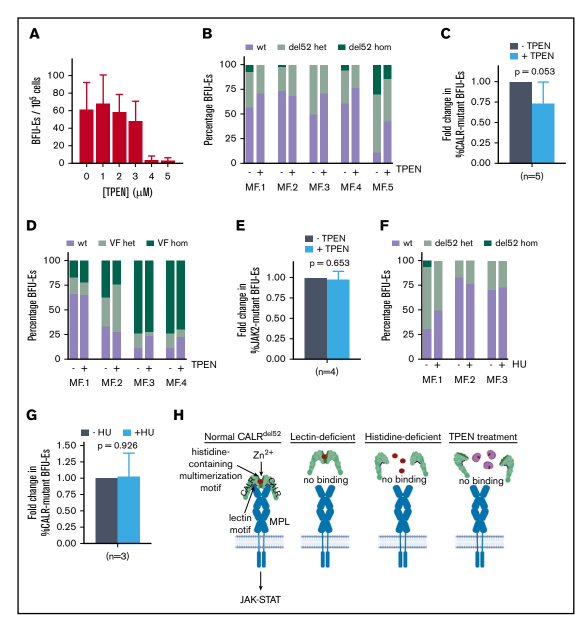


Figure 5. Zinc chelation exhibits selective cytotoxicity against CALR-mutant cells from MPN patients. (A) BFU-E colony counts of PBMCs from healthy volunteers grown in methylcellulose in the presence of 0 to 5 μ M TPEN. (B) PBMCs from 5 myelofibrosis patients (allele burden ranging from 20% to 73%) were cultured in the presence and absence of 2.5 μ M TPEN, and BFU-Es genotyped for the presence of CALR del52 mutation after 14 days. Proportion of colonies with wild-type CALR (wt), heterozygous for CALR del52 mutation (del52 het), or homozygous for CALR del52 mutation (del52 hom) cultured in the presence and absence of 2.5 μ M TPEN. At least 20 colonies were genotyped for each patient. (C) Average fold change of CALR-mutant colonies (including heterozygous and homozygous colonies) for 5 MF patients examined following TPEN treatment relative to untreated cultures, which was arbitrarily designated as 1. (D) Proportion of colonies with JAK2 (wt), heterozygous for JAK2V617F mutation (VF hom) cultured in the presence and absence of 2.5 μ M TPEN. At least 20 colonies of JAK2V617F mutation (VF hom) cultured in the presence and absence of 2.5 μ M TPEN. At least 20 colonies were genotyped for each patient. (E) Average fold change of JAK2-mutant colonies (including heterozygous and homozygous colonies) for 4 MF patients examined following TPEN treatment relative to untreated cultures, which was arbitrarily designated as 1. (F) Proportion of colonies with CALR (wt), heterozygous for CALR del52 mutation (del52 hom) cultured in the presence and absence of 2 μ M hydroxyurea (HU). At least 15 colonies were genotyped for each patient. (G) Average fold change of CALR-mutant colonies (including heterozygous and homozygous colonies) for 3 MF patients examined following HU treatment relative to untreated cultures, which was arbitrarily designated as 1. (H) Model depicting interplay between zinc, CALR^{del52} multimerization and MPL activation. (C-G) Testing for statistical significance was performed using Student *t* test (*

heavy metal chelators for treatment of *CALR*-mutated MPNs. Moreover, these data also open up the possibility that agents that interfere with CALR^{del52} multimerization could also be an effective therapeutic strategy for impairing CALR^{del52} oncogenic activity. In conclusion, our data provide several novel insights into the molecular mechanism by which CALR^{del52} interacts with MPL to induce MPN. Specifically, we (1) uncover a triad of zinc-binding histidines within CALR^{del52} required for its oncogenic activity; (2)

provide evidence that zinc-dependent homomultimerization of CALR^{del52} underlies its ability to bind MPL; and (3) determine that reduction of intracellular zinc abrogates the oncogenic activity of CALR^{del52} and could represent a new therapeutic modality in the treatment of *CALR*-mutant MPNs.

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Authorship

Contribution: J.F.R., A.J.B., D.G.K., A.M., and E.C. designed the research; J.F.R., A.J.B., F.N., G.B., R.S., H.P., E.L.B., G.A., S.A.B., B.R.J., P.L., A.M., and E.C. performed experiments and collected data; E.J.B. and A.R.G. provided patient samples; J.F.R., A.J.B., P.L., A.M., and E.C. analyzed the data; and J.F.R., A.J.B., F.N., A.M., and E.C. wrote the manuscript.

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