RESEARCH ARTICLE

JAK2 Exon 14 Skipping in Patients with Primary Myelofibrosis: A Minor Splice Variant Modulated by the JAK2-V617F Allele Burden

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Abstract

Background

Primary myelofibrosis (PMF) is an acquired clonal disease of the hematopoietic stem cell compartment, characterized by bone marrow fibrosis, anemia, splenomegaly and extramedullary hematopoiesis. About 60% of patients with PMF harbor a somatic mutation of the JAK2 gene (JAK2-V617F) in their hematopoietic lineage. Recently, a splicing isoform of JAK2, lacking exon 14 (JAK2Δ14) was described in patients affected by myeloproliferative diseases.

Materials and Methods

By using a specific RT-qPCR method, we measured the ratio between the splicing isoform and the JAK2 full-length transcript (JAK2+14) in granulocytes, isolated from peripheral blood, of forty-four patients with PMF and nine healthy donors.

Results

We found that JAK2Δ14 was only slightly increased in patients and, at variance with published data, the splicing isoform was also detectable in healthy controls. We also found that, in patients bearing the JAK2-V617F mutation, the percentage of mutated alleles correlated with the observed increase in JAK2Δ14. Homozygosity for the mutation was also associated with a higher level of JAK2+14. Bioinformatic analysis indicates the possibility that the G>T transversion may interfere with the correct splicing of exon 14 by modifying a splicing regulatory sequence.
Conclusions

Increased levels of \( JAK2 \) full-length transcript and a small but significant increase in \( JAK2 \) exon 14 skipping, are associated with the \( JAK2-V617F \) allele burden in PMF granulocytes. Our data do not confirm a previous claim that the production of the \( JAK2\Delta14 \) isoform is related to the pathogenesis of PMF.

Introduction

The human \( JAK2 \) gene occupies a genomic region of about 14 kilobases (kb) on the short arm of chromosome 9 (9p24.1); it produces a transcript of 5.3 kb consisting of 25 exons that is translated into a cytoplasmic tyrosine kinase of 1132 amino acids, and belongs to the Janus kinase family (NCBI gene ID: 3717). In myeloproliferative neoplasms (MPNs), a somatic guanine-thymine substitution (c.1849G>T) located in the terminal part of exon 14 of \( JAK2 \), has been identified [1–4]. The consequent amino acid change, valine 617 to phenylalanine (\( JAK2-V617F \)), alters the structure of the pseudokinase domain with important consequences in activation [5–7]. This mutation is observed in almost all patients with polycythemia vera and in more than half of those with essential thrombocythemia or primary myelofibrosis (PMF). The measure of the ratio between mutated and total alleles in genomic DNA extracted from granulocytes (usually designated as the \( JAK2-V617F \) allele burden) is used either at diagnosis for prognostic information or during treatment as a means to assess minimal residual disease [8].

By using the quantitative fragment length analysis technique, Ma et al. [9] described an alternative splicing event in the \( JAK2 \) gene, resulting in the missing exon 14 both in plasma and in granulocytes of patients with MPNs. The transcript was found in ratios ranging from 2% to 26% compared to the amount of the full-length isoform, and it was reported to be translated into a truncated protein of approximately 70 kDa. As it was detected only in patients with MPNs, and more likely in patients tested negative for \( JAK2-V617F \), it was suggested that the isoform could play a significant role in the pathophysiology of MPNs. The authors hypothesized that the truncated protein isoform dimerizes with the wild type \( JAK2 \), activating its kinase domain and consequently the \( JAK2-STAT \) pathway.

In this study, we assessed the exon 14-skipping variant (\( JAK2\Delta14 \)) in granulocytes of patients with PMF by using an isoform specific RT-qPCR method (S1, S2 Figs.) [10]. Moreover, we investigated the possible mechanism driving the alteration of splicing associated with the \( JAK2-V617F \) mutation.

Materials and Methods

Ethics statement

All work was performed according to a protocol approved by the Ethic Committee of the IRCCS Policlinico S. Matteo Foundation. Written informed consent was obtained from each patient before data were entered in the database.

Patients and samples

We tested peripheral blood samples of 44 patients with PMF selected from those referred to the Center for the Study of Myelofibrosis at the Fondazione IRCCS Policlinico S. Matteo. The diagnosis of PMF was based on 2008 WHO criteria [11]. Fourteen patients were \( JAK2-V617F \)
negative (wild type), and thirty positive for the V617F mutation (a cutoff of 1% mutated alleles was fixed as positive). In addition, we tested nine healthy control individuals.

The samples were collected using 0.105 M sodium citrate tubes (BD Vacutainer), stored at 4°C and processed within 4 hours after collection. Blood granulocytes were isolated from the lower interface of a Lympholyte-H density gradient (1.077 g/cm³ at 20°C, Cedarlane Laboratories Ltd.) and then submitted to erythrocyte lysis (BD Pharm Lyse, BD Biosciences).

Both DNA and RNA were extracted from granulocytes and cell lines (see subsequent paragraph "Cell line culture"). Total RNA was extracted with the miRNeasy Mini Kit (Qiagen) and further DNA purified by on-column digestion with the RNase-free DNase Set (Qiagen), according to the manufacturer’s instructions. Genomic DNA was extracted using the QIAamp DNA Blood Mini Kit (Qiagen). Nucleic acids were quantified with a Nanodrop 1000 spectrophotometer (Thermo Scientific).

cDNA synthesis was carried out using the iScript kit (Bio-Rad). In brief, 150 ng of each total RNA sample was reverse transcribed using a blend of oligo-dT and random primers, subsequently diluted with nuclease–free water to 3.75 ng/μL (total RNA equivalent) and stored at -80°C.

The quality of RNAs extracted from granulocytes and cell lines was assessed in two healthy individuals, four patients and one cell line, randomly chosen (S1 Table). The cDNAs resulting from reverse transcription of these RNA samples were analyzed, using qPCR methods, in order to test the following parameters: (i) RNA integrity, using a 50/30 ratio mRNA integrity assay [12] (qHsaCtlD0001002, Bio-Rad); (ii) contamination from genomic DNA, using an assay that targets an untranscribed region of the human genome (qHsaCtlD0001004, Bio-Rad); (iii) presence of PCR inhibitors, using a positive PCR control assay that targets a synthetic DNA added to the reaction mix (qHsaCtlD0001003, Bio-Rad).

Cell line culture

DAMI and K562 cell lines were cultured from laboratory stocks, while the UKE-1 cell line was generously provided by the original investigators [13]. Cells were routinely cultured in Iscove Modified Dulbecco’s Medium [IMDM] supplemented with 10% fetal bovine serum, 2% glutamine and 1% penicillin/streptomycin, at 37°C in a fully humidified incubator in the presence of 5% CO2. Where indicated, cycloheximide (CHX) (CAS 66-81-9, 25 mg/mL in DMSO) was added to the medium at a final concentration of 10 μg/mL, 8 hours before harvesting. Three independent experiments for each condition were performed using the same cell lines.

RT-qPCR gene expression analysis

Primers for EvaGreen assays (S2 Table) were designed using the Beacon Designer 7.9 software (Premier Biosoft International). Quantification of transcripts was carried out in a 15 μL reaction mix containing 1X SsoFast EvaGreen Supermix (Bio-Rad) and 400 nM of each primer. The PCR conditions were 95°C for 30 sec followed by 40 cycles of 95°C for 5 sec and 60°C for 5 sec. Melting curves were generated after amplification in the range 65–95°C with increments of 0.2°C every 10 sec. For each experiment, 3 μL of cDNA (corresponding to 11.3 ng of total RNA) were used. The PCR data were collected using the CFX96 Real-Time System (Bio-Rad). Each sample was tested in duplicate. Calculation of normalized relative expression levels was done using the Qbase Plus software version 2 (Biogazelle): a three-point serial dilution (1:4) of a mix of cDNA from patients and controls was included for each gene, to perform the amplification efficiencies correction; three samples were included in each run to generate an inter-run calibration; normalization was performed using the most stably expressed reference gene (or the two most stable, in the case of in vitro experiments) which was selected using the geNorm algorithm [14], with the following candidates: YWHAZ, GAPDH, HPRT1, UBC. Other authors
have validated, in nine human bone marrow samples, the expression stability of the above-mentioned reference genes [14].

**Standard curves**

The percentage of JAK2Δ14 compared to the full-length isoform JAK2+14 was calculated using absolute standard curves. The PCR products corresponding to the full-length transcript and skipped isoform (S2 Table) were run on 2% agarose gels in TBE buffer. The amplified fragments were excised and purified from the gel using QIAquick spin columns (Qiagen). The concentrations of the PCR products were measured using both the Quantifluor dsDNA System on a Quantifluor-ST fluorometer (Promega) and the Nanodrop 1000 spectrophotometer (Thermo Scientific). The molecular weight of the PCR fragments was determined using the software MacVector (MacVector, Inc.) and used for the conversion of micrograms to picomoles. Finally, equimolar dilutions of PCR fragments were used to generate the standard curves (S2 Fig.).

**JAK2-V617F allele burden in genomic DNA and cDNA**

JAK2-V617F allele burden in the genomic DNA and cDNA obtained by retrotranscription of total-RNA from granulocytes was measured by allele-specific quantitative PCR, as previously described [15]. The JAK2-V617F allele burden was calculated by comparison with a standard curve obtained by a dilution series of genomic DNA from a patient with 100% allele burden into donor wild type DNA, in the following proportions: 2%, 5%, 12%, 25%, 50%, 75%, 95%, and 100%. To quantify the number of mutated transcripts, a similar standard curve was obtained by mixing the cDNA from a 100% mutated patient into donor wild type cDNA with identical JAK2 expression levels. qPCR reactions were carried out in a 20 μL reaction mix containing 1X SsoFast Probes Supermix (Bio-Rad), 400 nM of each primer and 200 nM of 6-FAM/BHQ-1 hydrolysis probe (Sigma-Aldrich). The PCR conditions were 95°C for 2 min followed by 40 cycles of 95°C for 5 sec and 62°C for 10 sec. For each experiment, 45 ng of genomic DNA or 11.3 ng of cDNA, were used.

**Reverse Transcription-PCR (RT-PCR) experiments**

RT-PCR reactions (35 cycles, annealing temperature determined with a gradient between 50°C and 60°C, 1 min/kb extension) were performed in a C1000 thermal cycler (Bio-Rad) using granulocyte’s total RNA from patients and healthy individuals (see S3 Table for the list of the primers used). The amplifications were carried out using the GoTaq Flexi DNA Polymerase (Promega) according to the manufacturer’s instructions. The PCR products were run on a 2% agarose gel in TBE buffer and stained with 1x Sybr Gold (Invitrogen).

**Bioinformatic analysis**

The Human Splicing Finder 3.0 and ESEfinder 3.0 web servers were used to analyze the JAK2 exon 14 sequence to search for known exonic splicing enhancer and silencer (ESE/ESS) motifs, in wild type sequence and in the presence of the c.1849G>T (V617F) mutation. MacVector 12 (MacVector, Inc.) application was used for editing of JAK2 sequences (NG_009904.1, NM_004972.3), open reading frame analysis, protein prediction and calculation of the molecular weight of PCR products.

**Statistical analysis**

The relationship between measured variables was analyzed using bivariate linear regression and multiple regression analyses. The non-parametric Mann-Whitney U test was used to
compare the variables measured in patients and controls. Fisher’s PLSD post-hoc test and Student’s unpaired t-test were used to study, respectively, pairwise differences in gene expression between cell lines and normal granulocytes and between *in vitro* experimental conditions. Differences were declared significant below the significance level of 0.05. StatView version 5.0.1 (SAS Institute Inc.) was used for statistical analysis. Box plots were used to represent the distributions of variables. In box plots, horizontal lines display 25th, 50th, 75th percentiles and lowest and highest values of a variable. In scatter diagrams, dotted lines were used to represent the 95% confidence band of the regression line. Prism 4.0c (GraphPad Software Inc.) was used for data representation.

**Results**

**JAK2Δ14 in healthy controls and PMF patients**

Measurable levels of the spliced JAK2Δ14 variant were detected in blood granulocytes both in healthy controls and PMF patients. The median value of the JAK2Δ14 transcript, as expressed in percentage of the full-length transcript (JAK2+14), was higher in PMF patients than in controls without reaching statistical significance (0.71% vs. 0.42%, respectively; Mann-Whitney U test: *p* = 0.076).

However, while wild type patients had JAK2Δ14 levels that did not differ significantly from healthy donors (0.37% vs. 0.42%, respectively; Mann-Whitney *U* test: *p* = 0.378; Fig. 1), patients carrying the JAK2-V617F mutation showed significantly greater percentages of the spliced isoform with respect to controls (median: 1.0%, interquartile range: 0.49; Mann-Whitney *U* test: *p* < 0.01; Fig. 1). Moreover, in JAK2-V617F mutated patients, the level of the isoform was strictly correlated with the JAK2-V617F allele burden (*R*² = 0.43, *p* < 0.001; Fig. 2).

**Bioinformatic analysis of JAK2 exon 14**

We analyzed the possible effects of the c.1849G>T mutation on exonic splicing regulatory sequences (ESE or ESS) using ESEfinder 3.0 [16] and Human Splicing Finder 3.0 [17] programs. The JAK2 exon 14 consists of 88 base pairs and the mutation occurs at position 73. In the mutated sequence, two independent algorithms [16, 18] identified a possible alteration of an exonic ESE site. Another algorithm [19], designed to search for phylogenetically conserved sequences that can act as silencers or enhancers depending on exonic context, recognizes, in
the same sequence identified by ESEfinder (TGTGTC) a possible splicing regulatory element disrupted by the mutation (Fig. 3, S4 Table). The ESEfinder 3.0 analysis also showed that this sequence is a nearly optimal consensus motif for SRp55 (TGCGTC). Two other computational approaches [20, 21] predict the potential creation of an exonic splicing silencer (ESS). Conversely, in the same exonic subsequence, the ESEfinder algorithm recognizes the possible creation of an ESE sequence containing a SC35 binding motif (Fig. 3, S4 Table). Another matrice, implemented in Human Splice Finder 3.0, indicates hnRNP-A1 as a ligand of a potential ESS (TATGTG), but the software predicted only a slight enhancing effect of the G>T transversion on its function (S4 Table).

Regulation of JAK2 transcription

YWHAZ was used as a reference gene for expression studies in granulocytes because it was experimentally found to be the most stably expressed in these cells. In order to study the regulation of JAK2 gene transcription, we analyzed the level of expression of JAK2 full-length mRNA in patients with PMF and its relationship with the amount of the JAK2Δ14 splicing isoform. In agreement with previously reported data [22], the JAK2+14 transcript levels were significantly higher in patients with the highest V617F allele burden (Fig. 4). Indeed, we observed a median 50% increase of JAK2+14 in patients bearing the V617F mutation in more than 50% of alleles, compared to those with a wild type genotype (Mann-Whitney U test: p < 0.01).

Since the JAK2 exon 14 skipping, changes the open reading frame and results in the introduction of a premature termination codon (PTC) (S4 Fig.), we wondered whether JAK2Δ14 could be the target of the nonsense-mediated mRNA decay (NMD) system that is known to require the presence of a PTC at more than 50–55 nucleotides (nt) from the last junction between exons [23, 24]. With RT-PCR, we documented that the JAK2Δ14 transcript extends at least over exon 18 (S5 Fig., S3 Table).

The percentage of mutated transcripts in cDNA was measured to evaluate the hypothesis that a combination of NMD activity and preferential production of the isoform by pre-mRNA
containing the V617F mutation could lead to a decrease in production of JAK2+14 mutated transcripts in the samples positive for the JAK2-V617F mutation. Conversely, in agreement with another study [15], we observed that the proportion of JAK2-V617F mutated alleles, was the same for both genomic DNA and cDNA (Fig. 5).

JAK2Δ14 in cell lines bearing the JAK2-V617F mutation

In order to assess the effect of the JAK2-V617F mutation on JAK2 exon 14 skipping in cells other than granulocytes, we assayed the expression of JAK2 main transcript (JAK2+14) and the relative level of JAK2Δ14 in cell lines either JAK2-V617F homozygous (UKE-1, DAMI) or wild type (K562) [25]. In K562 and UKE-1 lines, the expression of JAK2+14 was lower than that observed in normal granulocytes while in DAMI, the presence of many copies of the gene [25] caused mRNA levels that were more than two times higher than in normal granulocytes. Nevertheless, the relative amount of JAK2Δ14 in all three cell lines was lower than that measured in granulocytes: between 20 and 40% of the average value observed in granulocytes (Fig. 6a).
The absence of an enhancing effect of the c.1849G>T mutation on the level of the exon 14-skipping isoform in the JAK2-V617F homozygous cell lines could be due to several factors. We tested the hypotheses that different concentrations of splicing factors in these cells and/or a higher degradation due to the NMD system may maintain JAK2Δ14 at low levels. To assess the first hypothesis, we measured the mRNA levels of two splicing factors indicated in bioinformatics analysis: SRp55 and hnRNP-A1. In all three cell lines, the levels of both mRNAs were vastly higher than those observed in granulocytes: about 10 times for SRp55 and between 26 and 50 times for hnRNP-A1 (Fig. 6b). To investigate the possibility of NMD system
involvement, we treated the above-mentioned cell lines with CHX, a protein synthesis inhibitor that locks the NMD system activity [24]. To verify the effectiveness of the treatment we measured the expression of a SRp55 splicing variant containing a PTC (SRp55-PTC+b). It has been demonstrated that the inhibition of the NMD system, both with CHX and through depletion of UPF1, causes an increase of this variant in HeLa cells [26]. Our experiment confirms the results obtained by Lareau et al. [26]. Eight hours after treatment, we observed a significant increase in the levels of SRp55-PTC+b messenger in all cell lines. On the contrary, neither the level of JAK2+14 nor that of JAK2Δ14, were significantly changed after treatment with CHX (S3 Fig.).

Discussion

Besides affecting the amino acid sequence, which in turn is critical for the function of the protein, missense and nonsense mutations can also alter splicing regulatory sequences, that lead to an incorrectly spliced transcript [27, 28].

With this study we characterized an exon 14-skipping isoform of the JAK2 gene that is mutated in approximately 60% of patients with PMF. We found that JAK2 exon 14 skipping occurs constitutively both in healthy individuals and PMF patients. In PMF patients bearing the JAK2-V617F mutation, the production of the skipped isoform correlated with the percentage of mutated alleles. This observation, combined with the results of bioinformatic analysis of the JAK2 exon 14 sequence, allowed us to hypothesize that the c.1849G>T somatic transversion, in addition to determining the amino acid substitution p.V617F, could change a splicing regulatory sequence, causing an increase in the production of the skipping isoform in mutated subjects. However, even in the presence of high JAK2-V617F allele burden, the amount of isoform represented no more than 2.5 percent of the full-length transcript. Therefore, having found some evidence that JAK2Δ14 could meet the criteria as the target of NMD, we asked whether this system intervenes by degrading the isoform and consequently, minimizing the potential...
damage due to a hypothetical abundant production of JAK2Δ14 caused by the JAK2-V617F mutation.

As a matter of fact, in-frame nonsense codons located upstream of the last junction between exons were recognized as PTCs and targeted the mRNA for degradation. Nevertheless, a study by Pan et al. [29] showed that the majority of transcripts containing PTCs generated by alternative splicing (which accounts for about one third of the splice variants), are present at low levels, and that only a small fraction of these is regulated by the NMD system. It is not clear to what extent such variants are functionally relevant, but a recent deep sequencing analysis of the human lymphoblastoid cell transcriptome [30] seemed to confirm the hypothesis that a large fraction may arise as a consequence of the probabilistic nature of the splice sites recognition, and can be classified as non-functional “noise” [31]. Based on the above-mentioned results and on the analysis of the percentage of the c.1849G>T mutated alleles in cDNA compared to genomic DNA, we infer that the overproduction of the isoform might be minimal. The absence of a significant effect of the increased production of JAK2Δ14 on the expression of the mutated alleles, led us to conclude that the observed low level of this splice variant was probably due to its limited production rather than to a massive degradation operated by the NMD system. Indeed, we could not detect any significant enhancement in the levels of JAK2Δ14 following NMD inhibition with CHX in model cell lines.

In order to explain why the presence of a homozygous mutation does not affect the production of JAK2Δ14 in DAMI and UKE-1 cells, we proposed that a different concentration of splicing factors in these cell lines could maintain JAK2Δ14 at low levels. Indeed, the transcript levels of hnRNP-A1 and SRp55 (i.e. two splicing factor suggested by the bioinformatics analysis) are one order of magnitude higher in cell lines compared to their expression levels in granulocytes. Previous studies showed that altered concentrations of SRp55 and hnRNP-A1 determine quantitative changes in the ratio between isoforms of cancer related genes [32–34]. Assuming that the levels of hnRNP-A1 and SRp55 mRNAs are directly proportional to the amount of the related splicing factors, we tested two alternative hypotheses. The first was that the mutation increases the binding capacity of an ESS bound by the factor hnRNP-A1. This conjecture is contradicted by what was observed in the cell lines above, where the high levels of hnRNP-A1 should lead to higher activity of the linked ESS and a consequential increase in JAK2Δ14 levels. The second was that the mutation disrupts an ESE linked by the SRp55 protein. This hypothesis is compatible with our observations because the high levels of SRp55 in DAMI and UKE-1 cells could compensate for the predicted interference caused by the c.1849G>T mutation with the binding of this factor. These findings together with the above-discussed results, although not sufficient to derive definitive conclusions, support the initial hypothesis that the mutation interferes with the splicing of exon 14 through the modification of a splicing regulatory sequence. Further experiments are needed to confirm this hypothesis and to analyze the different possibilities that emerged from computational analysis.

Another result of this study is that the JAK2-V617F mutation was also associated with an increased amount of full-length isoform JAK2+14. Also in this case, the effect was proportional to the percentage of mutated alleles and in homozygous patients consisted in an average 50% increase of JAK2+14 levels. Although our data do not allow clarification of the mechanism that determines the increase in transcript levels, this observation may support a previously proposed hypothesis raised to explain why the individuals carrying the 46/1 haplotype have an increased risk of developing the mutation [35]. In accordance with the “fertile ground” hypothesis [36], the mutation occurs with the same probability on the different alleles, but the cells in which the mutation occurs on 46/1 haplotype have a selective advantage. It can be hypothesized that the observed increment in JAK2 mRNA levels is caused by the occurrence of
the JAK2-V617F mutation on the 46/1 haplotype. In this case, the increased production of the mutant JAK2 protein could contribute to the above-mentioned selective advantage.

Our approach did not confirm the presence of high amounts of JAK2Δ14 observed by Ma et al. [9]. This could be due to the fact that the Quantitative Fragment Length Analysis technique, originally developed for the prenatal diagnosis of chromosomal abnormalities [37], used by Ma et al., is less suitable for the quantification of splice variants. Since with this method, fragments of different sizes are simultaneously amplified, overestimation of the amount of the isoform that produces a shorter fragment is possible because it tends to be amplified preferentially with respect to the full-length counterpart. Moreover, if the amplification is not limited to the exponential phase, the least represented isoform is overestimated [37–39].

The experimental evidence described here argues against the hypothesis that the presence of this splice variant could be pathogenetically associated with MPNs. It is unlikely that the slight increase in the amount of JAK2Δ14 could produce a truncated protein at significant levels. In addition, the fact that the level of JAK2Δ14 is comparable in healthy subjects and in patients is in contrast with the hypothesis that its presence could be involved in the pathogenesis of PMF. Moreover, it was observed that the ectopic expression of a truncated protein isoform of JAK2 lacking the protein kinase domain (Jak2-829), has the effect of blocking the erythropoietin-dependent inhibition of apoptosis [40]. It can be hypothesized from the above observation that the production of a truncated protein isoform of JAK2, resulting from translation of JAK2Δ14, could have an antiproliferative effect that would be desirable in MPNs.

Supporting Information
S1 Fig. JAK2Δ14 RT-qPCR analysis in healthy controls and PMF patients. EvaGreen amplification signals for YWHAZ, JAK2+14 and JAK2Δ14, in two individuals with normal (grey) and increased (black) level of the exon 14-skipping isoform. Top left box shows melting peaks obtained by High Resolution Melting Analysis of the three amplification products: it can be observed the different melting peak morphology caused by the JAK2-V617F mutation present in the JAK2+14 transcripts of the patient with increased level of JAK2Δ14. (JPG)

S2 Fig. Quantification of PCR-JAK2+14 and PCR-JAK2Δ14 by absolute standard curves. Equimolar dilutions of PCR-JAK2Δ14 (left) and PCR-JAK2+14 (right) amplicons, were used to generate two standard curves utilized to calculate the percentage of alternative transcript. The three points correspond to 1:4 serial dilutions of the gel-purified PCR products. (JPG)

S3 Fig. Effect of CHX treatment on JAK2 alternative transcripts containing PTCs. RT-qPCR was used to assay mRNAs levels in cell lines either homozygous for the JAK2-V617F mutation (UKE-1, DAMI) or wild type (K562). Transcript level ratios between CHX-treated (orange) and untreated cells (blue), are shown for: SRp55 constitutive transcript (SRp55), SRp55 PTC-containing isomorf (SRp55-PTC+b), JAK2 full-length transcript (JAK2+14) and JAK2 exon 14 skipping isomorf (JAK2Δ14). Data are expressed as means (± SEM) of three independent experiments performed using the same cell line. Normalized expression of targets genes was obtained using the two genes with the lowest geNorm M-value: YWHAZ/HPRT1 for DAMI, GAPDH/HPRT1 for K562 and UKE-1. Asterisks (*) indicate significant changes in gene expression after treatment. (JPG)

S4 Fig. Hypothetical translations of the JAK2Δ14 subsequence resulting from the junction between exons 13 and 15. The sense strand (black), its complementary strand (blue) and
their possible phases of translation, are shown. Single-letter code is used to represent the amino acids. A stop codon is indicated by an asterisk (\(^*\)). The reading frame, used in the translation of the full-length transcript (JAK2+14), is represented in the first row above the sense strand.

S5 Fig. The alternative transcript (JAK2Δ14) extends at least until exon 18 and can be the target of the Nonsense Mediated Decay (NMD) system. (A) The diagram shows the location of the primers in the JAK2 full-length mRNA (above) and in the isoform lacking exon 14 (below). As in the qPCR, forward primers were specific for each isoform while the reverse primer was, in both amplifications, localized in exon 18 (S2 Table). In the alternative isoform, the hypothetical position of the stop codon and exon junction complexes (which is expected to activate the NMD system), are indicated. (B) Electrophoresis of PCR products obtained by amplifying the cDNA of a patient with 2.5% level of JAK2Δ14 isoform, at three different annealing temperatures. The expected amplicon sizes are 495 bp for the JAK2Δ14 isoform (PCR-Δex14/ex18) and 556 bp for the JAK2+14 constitutive isoform (PCR-ex14/ex18).

S1 Table. RNA quality control assay results for seven randomly chosen cDNA samples used in this study.

S2 Table. Primers used in RT-qPCR experiments.

S3 Table. Primers used in RT-PCR experiments.

S4 Table. Identification of potential splicing regulatory sequences in wild type and V617F mutated JAK2 exon 14 sequences. Twelve different matrices implemented in Human Splicing Finder (HSF) 3.0, are presented.

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Author Contributions

Conceived and designed the experiments: PC GB VR. Performed the experiments: PC. Analyzed the data: PC. Contributed reagents/materials/analysis tools: VP LV RB MP MZ. Wrote the paper: PC VR GM GB.
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