TLR8-dependent TNF-α overexpression in Fanconi anemia group C cells

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Introduction

The Fanconi anemia (FA) proteins play an important role in regulating genome stability,1 but there is little evidence that the loss of the genoprotection per se in FA cells accounts for the molecular pathogenesis of the bone-marrow failure characteristic of this disease. In fact there is evidence that at least some of these proteins are multifunctional2 and participate in canonical signaling pathways in hematopoietic cells.2–8 Fanconi anemia, complementation group C (FANCC)–deficient cells, for example, are hypersensitive to the apoptotic effects of tumor necrosis factor-α (TNF-α).4–9 In addition, FA cells overproduce TNF-α for reasons that have not yet been fully explained.10–12 Most importantly, there is clear evidence that overproduction of and hypersensitivity to TNF-α in hematopoietic cells of FANCC−/− mice results in bone marrow hypoplasia13,14 and that long-term ex vivo exposure of murine FANCC−/− hematopoietic cells to both growth factors and TNF-α results in the evolution of cytogenetically marked preleukemic clones.9 Therefore, the hematopoietic phenotype of FA may evolve from the overproduction of precisely the cytokine to which FA stem cells are hypersensitive. We designed gene expression microarray experiments by using marrow cells from both patients with FA and normal volunteers in part to seek potential clues to the mechanisms by which FA cells overproduce TNF-α.

Recognizing that transcriptional analysis would not reveal aspects of the FA phenotype that were controlled translationally or posttranslationally, we also conducted a proteomics analysis. We based our experimental design on an accepted function of the FA “nuclear core complex,” that is, its capacity to facilitate monoubiquitinylation of both Fanconi anemia, complementation group I and Fanconi anemia, complementation group D2 (FANCD2).15,16 Although it is clear that monoubiquitinylation, at least of FANCD2, is required for the avoidance of genotoxicity,17 it seemed to us unlikely that 8 individual FA genes encoding the “core complex proteins” should have evolved to control the monoubiquitinylation of merely 1 or 2 nuclear proteins. Therefore, reasoning that ubiquitinylation of a variety of other proteins might also be influenced by the core FA proteins, we designed a proteomics survey of ubiquitinylated proteins in FA-C cells and isogenic controls. We reasoned that this approach might lead to the identification of other proteins underubiquitinylated in mutant cells. As reported herein, the gene expression microarray analysis revealed a significant overrepresentation of overexpressed ubiquitin pathway genes in the mutant cells. We therefore took into account the alternative possibility that some ubiquitinylated proteins might be found uniquely in the mutant cells.

Indeed, one such protein, Toll-like receptor 8 (TLR8), did appear in the ubiquitin-positive fractions only in FANCC-mutant cells. Given that TLR8 activation is known to induce expression of TNF-α, we focused our postproteomics functional studies specifically on the TLR8 pathway. The results of our studies demonstrate that (1) FANCC modulates the activation state of TLR8 by...
suppressing either its ubiquitylation or its association with another ubiquitylated protein; (2) that FANCC inactivation results in excessive TNF-α gene expression that results specifically from the inappropriate activation of TLR8; and (3) that this function of the FANCC protein has structural requirements that the canonical genome protective function of FANCC does not.

Methods

Gene expression microarray analysis

Patients. Aspirated bone-marrow samples from patients with FA were obtained in 1 of 2 centers: Oregon Health & Science University, Portland, or Hospital de Clinicas, Federal University of Parana, Curitiba, Brazil. Patients with FA who were eligible for this study met the following 3 on-study criteria: (1) a positive chromosome breakage test on exposure of either lymphoblasts or fibroblasts to either diepoxybutane or MMC or both; (2) a normal bone marrow cytogenetics study obtained by the use of conventional Giemsa banding methods on metaphase preparations within 12 months of accrual to the study; and (3) absence of acute myelogenous leukemia. All patients with clonal cytogenetic abnormalities and all who had received a stem cell transplant were ineligible. Bone marrow samples were aspirated in heparinized syringes from 11 normal volunteers and 22 patients with FA meeting the on-study criteria. All human studies were approved by the institutional review boards of all participating institutions, and samples were obtained with informed consent in accordance with the Declaration of Helsinki.

RNA isolation and processing. Low-density marrow mononuclear cells were prepared from heparinized bone-marrow aspirates by the use of Ficoll-Paque. RNA prepared in Brazil was prepared immediately, frozen at −80°C (in water), and shipped on dry ice to Portland. Total RNA was extracted in both centers by use of the RNAsena Mini kit (Qiagen). Before hybridization to a GeneChip Human Genome HG-U133A (22,283 probe sets) arrays. Detailed methods and quality control measures are reviewed in the supplemental Methods (available on the Blood website; see the Supplemental Materials link at the top of the online article). Data visualization and exploratory analysis, including principal component analysis and hierarchical clustering, were performed by the use of Partek Genomics Suite (Partek) and GeneSifter (Geospiza). CEL files and probe set signals have been deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO Series accession number GSE16334 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE16334).

Murine splenocytes. Facc−/− mice were generated as described previously.9 Splenocytes from both wild-type and Facc−/− mice were isolated by first preparing single-cell suspensions by passing the spleens through a 40-μm mesh (Sigma-Aldrich) and then isolating the mononuclear cells by the use of Ficoll-Paque Plus (GE Healthcare). CD11b− cells were isolated by the use of antibody-conjugated magnetic beads (Easy Sep; StemCell Technologies). Cells were then cultured at a concentration of 2000 cells/200 μL for 24 hours in RPMI and 10% fetal calf serum (HyClone) in the presence of multiple doses of R848 (Alesxe Biochemicals). Supernatants were harvested for TNF-α quantification by enzyme-linked immunosorbent assay (ELISA; R&D Systems). All murine studies were approved by the Portland VA Institutional Animal Care and Use Committee.

Antibodies and reagents. The following rabbit polyclonal antibodies were purchased from Cell Signaling Technology: anti-phospho-IskB kinase (IKK)-α/β (Ser176/180), anti-IKK-α, anti-IKK-β, anti–phospho-interleukin-1 receptor–associated kinase (IRAK)–1 (Ser376), and anti–IRAK-1. Other rabbit polyclonal antibodies used include anti-ubiquitin (Santa Cruz Biotechnology) and anti–TLR8 (Abcam). Anti-FANC D2 was also purchased from Santa Cruz Biotechnology. The anti–TLR8 monoclonal antibody was obtained from MBL International. Actinomycin D (Sigma-Aldrich) was used at 5 μg/mL. The IRAK-1/4 inhibitor N-(2-morpholinoethyl)-2-(3-nitrobenzoylamido)-benzimidazole (Sigma-Aldrich) was used at 25 μM or 50 μM as indicated. All TLR ligands except R-848 were obtained from InvivoGen. R-848 was purchased from Alexis Biochemicals.

Cell lines. Epstein-Barr virus (EBV)–transformed lymphoblast cell lines HSC536N (FA-C) and PD149 (FA-C), and the corrected counterparts HSC536N/FANCC (FA-C/C) and PD149/FANCC (FA-C/C) were previously described.10,20 HSC536N (FA-C) cells expressing FANCC with site-directed mutations (F64A, T66A, S249A, E251A, F525A, and Y531A) were described previously.2 THP-1 cells were obtained from ATCC and THP-1 Blue cells were obtained from InvivoGen. All cell lines were grown in RPMI 1640 supplemented with 15% fetal bovine serum, 1% glutamine, 100 units/mL penicillin, 100 mg/mL streptomycin at 37°C, and 5% CO2 in a humidified atmosphere.

In vitro (His)6-ubiquitin conjugation and affinity chromatography. The proteomics-based method used to isolate ubiquitylated proteins is modified from a published procedure developed by one of the authors (D.A.)21 Specifically, hexahistidine-tagged ubiquitin, an adenosine triphosphate (ATP) recycling system, and inhibitors of the proteasome and deubiquitylating enzymes, were added to lysed FA cells and normal cells. This method uses endogenous enzymes systems (E1 ubiquitin–activating enzyme, E2 ubiquitin–conjugating enzymes, and E3 ubiquitin–protein ligases) present in the cell lysates. After the ubiquitylation reaction, ubiquitylated proteins were affinity purified by nickel chromatography, digested, and analyzed by 2D capillary LC-MS/MS. Subconfluent HSC536N and HSC536N/FA-C cells were washed twice with ice-cold Dulbecco phosphate-buffered saline (Gibco). Cells were then lysed in buffer containing 50mM Tris-HCl, pH 7.4; 0.15M NaCl; 1% Triton X-100; 1mM dithiothreitol; 2mM sodium orthovanadate; 1mM phenylmethysulfonyl fluoride; 1% leupeptin; 1% pepstatin; and 1% aprotinin. Cell extracts were centrifuged for 15 minutes at 4°C. Supernatants (11 mg) were incubated for 2 hours at 20°C with 1 mg of hexahistidine-tagged ubiquitin (BostonBiochem), 5μM ubiquitin aldehyde (BostonBiochem), and 10μM MG132 (BostonBiochem) in a final volume of 2.5 mL. Reactions were performed with and without energy regeneration system (BostonBiochem), which contains MgCl2, ATP, and ATP-regenerating enzymes. The samples were then desalted with PD10 desalting columns (Amersham Biosciences) and eluted in buffer containing 50mM Tris-HCl, pH 7.4; 0.15M NaCl; 1% Triton X-100; 300mM NaCl; and 10mM imidazole. The eluates were loaded onto HisTrap HP columns (Amersham Biosciences) and washed with buffer containing 50mM Tris-HCl, pH 7.4; 0.15M NaCl; 1% Triton X-100; 300mM NaCl; and 40mM imidazole.

Resin containing the bound His-tagged proteins was then removed from the columns by centrifugation and proteolytically digested with Lys-C endoproteinase and modified trypsin as described previously.21 The micro-capillary 2D LC-MS/MS methods were performed as previously described.21 In brief, after heat-denaturation, reduction of disulfide bonds and alkylation of cysteines with iodoacetamide, the digested samples were injected into a biphasic microcapillary high-performance liquid chromatography column and separated by cation-exchange in the first dimension and by reversed phase in the second dimension. Mass spectra were obtained on an LTQ-FT hybrid linear ion trap Fourier transform ion cyclotron resonance mass spectrometer (ThermoFinnigan), and peptide precursor-MS/MS spectrum pairs were analyzed by the use of SEQUEST and support vector machine learning.23

Immunoprecipitations. Total proteins were extracted on ice in buffer containing 50mM Tris-HCl, pH 7.4; 0.15M NaCl; 1% Triton X-100; 2mM sodium orthovanadate; 1mM phenylmethysulfonyl fluoride; 1% leupeptin; 1% pepstatin; and 1% aprotinin. Cell extracts were centrifuged for 15 minutes at 4°C to remove cellular debris. For immunoprecipitations, supernatants (1 ng of total protein) were preclarified by adding 30 μL (50% vol/vol) of Sepharose and incubated for 45 minutes at 4°C. The extracts were then incubated with indicated antibodies overnight at 4°C. The immune complexes were then bound to protein A (or G) Sepharose (1 hour, 4°C) and washed 2 times with lysis buffer, followed by one wash with phosphate-buffered saline. Samples were eluted in 15 μL of 2X Laemmli buffer (1M Tris-HCl, pH 6.8; 4% sodium dodecyl sulfate; 40% glycerol; and 4% 2-mercaptoethanol).
Immunooassays and immunoblotting. TNF-α levels in culture supernatants were measured using ELISA kits from R&D Systems according to the manufacturer’s protocol. Proteins were separated in sodium dodecyl sulfate–polyacrylamide gel electrophoresis gels under reducing conditions. The proteins were subsequently transferred to nitrocellulose and blocked with 5% nonfat milk (Nestle USA Inc) in Tris-buffered saline Tween-20 (0.1M Tris–HCl; 0.15M NaCl, pH 7.4; 0.05% Tween-20). Blots were incubated with indicated antibodies in 1% milk overnight at 4°C. Five percent bovine serum albumin was used rather than 1% milk in incubations with phosho-specific antibodies. After incubations with appropriate horseradish peroxidase-coupled secondary antibodies (Bio-Rad), proteins were detected by use of the Enhanced Chemiluminescence Kit (Amersham Biosciences).

Real-time reverse-transcription polymerase chain reaction. Total RNA was prepared from 1 to 5 × 10⁶ cells by use of the RNeasy Mini kit (QIAGEN). Complementary DNA synthesis and real-time polymerase chain reaction (PCR) were performed as described previously. Pre-designed primer and probe sets for TNF-α were purchased as Taqman Gene Expression Assays (Applied Biosystems [Hs00174128_m1]).

Electromobility shift assay. Nuclear extracts were prepared and binding reactions carried out as described previously. The sequence of the oligonucleotide used, containing the human interferon regulatory factor-1 nuclear factor (NF)-κB binding site, was 5′-CGG GCC GGG GAA TCC CGC TAA G-3′. The oligonucleotide was synthesized in at the Molecular Biology Core Lab (Portland VAMC) and labeled with [γ³²P]-ATP by the use of T4 kinase (Roche Applied Science).

siRNA. Cells were transfected by use of the Amaza nucleofection technology (Amaza Inc). siRNA against FANCC (SMARTpool) and control siRNA were purchased from Dharmacon. Following Amaza protocols, 4 × 10⁶ cells were suspended in 0.1 mL of solution V, mixed with 200 pmol of siRNA, and transfected by use of program A-23 of the Nucleofector device. After transfection, cells were immediately transferred to 0.5 mL of RPMI medium without serum in 6-well plates at 37°C. After 15 minutes’ incubation, 1.5 mL of complete medium was added, and the cells were cultured for 72 hours.

Stable suppression of FANCC with shRNA lentivirus. Lentiviral particles targeting FANCC and TLR8 and control (nontarget) shRNA was purchased from Sigma-Aldrich. The target and shRNA sequences are shown in supplemental Table 4. Transductions into THP-1 and THP-1 Blue cells were performed as described by the manufacturer. The media were replaced 1 day after transduction. After 2 more days, shRNA-expressing cells were selected with 0.6 µg/mL puromycin for 2 to 3 weeks, at which time the cells were maintained in media containing 0.3 µg/mL puromycin. For dual suppression of FANCC and TLR8, FANCC shRNA was expressed by use of the vector pLKO.1-hPGK-Neo-CMV-tGFP, which allowed selection of stable FANCC shRNA-expressing cells using neomycin (200 µg/mL).

NF-κB reporter assay. THP-1 Blue cells express secreted embryonic alkaline phosphatase (SEAP) under the control of a promoter inducible by NFκB. SEAP expression is maintained by growing the cells in media containing zeocin (InvivoGen). Upon NF-κB activation, the expression and secretion of SEAP is monitored by the colorimetric enzyme assay QUANTI-Blue (InvivoGen), which produces a purple-blue color that is measurable on a spectrophotometer at 650 nm.

A 2-way comparison of bone marrow RNA from 11 normal volunteers and 21 FA patients yielded a total of 2204 genes expressed differentially (1.5-fold change). Ontologic analysis was conducted by the use of GeneSifter on genes differentially expressed in FA bone marrow cells (FAapl) compared with normal marrow cells (NC). From the list of 1952 genes suppressed or overexpressed in FAapl samples by 1.5-fold or greater (P < .01, adjusted for false-discovery rate by use of the Benjamini and Hochberg method), the ontologic categories: “protein ubiquitination” (z = 6.71), “ubiquitin-dependent protein catabolic process” (z = 6.64), “regulation of ubiquitin protein ligase activity during mitotic cell cycle” (z = 4.24), and “negative regulation of ubiquitin protein ligase activity” (z = 3.79) were significantly overrepresented, as was the expected category of “negative regulation of programmed cell death” (z = 5.08; supplemental Figure 2). Moreover, the 2 highest-ranked (by z score) categories applied specifically to genes overexpressed in FA cells. Overrepresentation of ubiquitin related ontologies was not peculiar to FANCC RNA samples and persisted even when subsets of the FA samples were analyzed (FANCA alone, FANCC alone, and both FANCC and Fanconi anemia, complementation group G; not shown). The sample sizes do not permit us to determine whether this ontologic overrepresentation is similar across the 3 complementation groups and allows us to draw no conclusions regarding the 10 complementation groups not known to be represented in our cohort.

Differential protein ubiquitinylation in FA-C cells

Initially we designed our proteomic analysis of the ubiquitome in FA cells because we expected that FA cells would contain fewer ubiquitinylated proteins than complemented cells. However, the transcriptional observations (in which some ubiquitin-related genes were overexpressed in the FA group) suggested that FA cells might exhibit enhanced activity of some ubiquitinylation pathways as well. We performed in vitro ubiquitinylation reactions by using hexahistidine-tagged ubiquitin. All the necessary endogenous enzyme systems (E1, E2, and E3) were present in the cell lysates, and because this is an ATP-dependent process, ATP and ATP-regenerating enzymes were included. False positives, identified in samples in which ATP and ATP-regenerating enzymes were not included, were removed from our lists. Ninety-nine proteins were uniquely ubiquitinylated in the FA-complemented (FA-C/C) cell lysate but not the FA-C cell lysate (supplemental Table 3). On this list, the prevalence of proteins known to be ubiquitinylated directly ubiquitinylated by mass spectrometry.

Of relevance to the work described herein, we also identified 90 proteins that were ubiquitinylated in the Fanconi anemia cell lysate but not in lysates of complemented cells (supplemental Table 2). TLR8 was one of these. The TLR8 peptide sequences identified in supplemental Figure 2. Three other peptides of potential interest included IKKβ (supplemental Table 2), BRCA2 (supplemental Table 2), and SH3BP5 (supplemental Table 3). We used communoprecipitation methods (antib ubiquitin antibodies and antibodies targeting these 3 proteins) in an attempt to confirm the proteomics result, but these studies were negative. We attribute the negative results to the insensitivity of the communoprecipitation method in light of the unambiguous observation that SH3BP5 was directly ubiquitinylated by mass spectrometry.

Results

Dysregulation of ubiquitin pathway–related genes in FA-C mutant cells

The representation of specific complementation groups is outlined in supplemental Table 1. The FA group included 9 with mutations of Fanconi anemia, complementation group A (FANCA), 4 with mutations of FANCC, 4 with mutations of Fanconi anemia, complementation group G, and 5 with unassigned complementation groups.
FANCC deficiency influences TLR8 but not other TLRs

Because Fanc−/− mice are hypersensitive to lipopolysaccharide (LPS),13,14 we hypothesized that FA cells might be generally hyperresponsive to a variety of TLR agonists. The human monocytic cell line THP-1 Blue, which expresses an NF-κB-inducible reporter, SEAP, was transfected with 5 lentiviral shRNAs targeting FANCC (Sigma-Aldrich), one of which potently suppressed FANCC and suppressed MMC-induced FANCD2 mono-ubiquitination (supplemental Figure 4). In addition, compared with untransduced cells or cells transfected with nontargeted shRNA, the FANCC shRNA-transduced cells showed greater chromosomal instability in 2 separate experiments. Specifically, when exposed to MMC (80 ng/mL) for 3 days, there was an increase in the number of cells exhibiting at least one chromosomal break (36/50 shFANCC, 24/47 untransduced control cells) and an increase in the number of quadriradial forms (22/50 shFANCC, 10/50 untransduced control cells). No chromosomal breakage differences were observed when nontargeted shRNA transfected cells and the untransduced cells were compared in the same experiment (not shown). Cells were incubated with multiple doses of the TLR8 inducers CL075 and R848 for 24 hours after which SEAP was quantified colorimetrically. Both R848 (Figure 3) and CL075 (not shown) induced significantly greater levels of TNF-α and SEAP production in cells expressing FANCC shRNA compared with either untransduced cells or cells transfected with control shRNA (nontargeted). No other TLR ligands, including the TLR7-specific agonist imiquimod, induced a differential response in cells bearing FANCC shRNA (supplemental Figure 5).

The use of THP1 cells permitted us to confirm the involvement of TLR8, IKK, and IRAK in functional activation of NF-κB. Specifically, R848-induced activation of NF-κB in FANCC-deficient THP-1 blue cells was markedly suppressed by IKK and IRAK inhibitors and shRNA targeting TLR8 (Figure 4). Because of the role played by p38 mitogen-activated protein kinase (MAPK) in the TLR pathway, we also tested the hypothesis that TLR8-dependent TNF-α production depends upon p38 activation. As shown in Figure 4E-G, the p38 MAPK inhibitor SB203580 suppressed R848 induced NF-κB activation (Figure 4E) and TNF-α expression (Figure 4F) and also inhibited ground-state TNF-α production in FANCC-deficient lymphoblasts (Figure 4G).

It should be mentioned here that neither of the 2 FA-C patient cell lines (EBV-transformed lymphoblasts) responded to R848. Neither complemented nor mutant lymphoblasts exhibited enhanced TNF-α production after exposure to that ligand. We also found that splenic lymphocytes were unresponsive to LPS and R848 but that splenic macrophages were responsive to both (not shown) and were differentially responsive. That is, macrophages obtained from the spleens of Fanc−/− knockout mice produced more TNF-α in response to R848 than did macrophages from wild-type mice.

Lack of differential FANCC and FANCA interactions with TLR8 signaling factors

By using coimmunoprecipitation methods, we examined the interaction of 2 FA proteins, FANCA and FANCC, with TLR8 and its signaling intermediates in lymphoblasts and although we found associations between (1) FANCA and TRAF6 and (2) FANCA and TAK1 (not shown), these interactions were equivalent in FA-C cells and complemented cells. FANCC did not associate with A20, an inhibitor of both TLR and TNF receptor signaling pathways (not shown). There was no detectable FANCC/TLR8 interaction.
complex in either complemented or mutant FA-C lymphoblasts. By use of coimmunoprecipitation, we found there were no detectable FANCC or FANCA interactions with other proteins in the TLR pathway, including Myd88, IRAK-1, IRAK-4, IKK-α, IKK-β, NEMO, IκBα, IκBβ, IκBγ, Triad3A, suppressor of cytokine signaling 1, and IRAK-M (not shown).

Structure-function analysis indicates that several conserved regions of FANCC are required for normal TNF-α production

Although the precise biochemical mechanism for FANCC’s effect on TNF-α production remains unknown, we asked whether several conserved regions of FANCC were required for normal modulation of TNF-α. Six alanine substitutions of FANCC cDNA were created, 2 each in 3 highly conserved domains (Figure 5); these mutations are known to complement FA-C cells in the MMC sensitivity assay. These mutants as well as normal FANCC cDNA and a disease-causing truncation mutant cDNA (M55) were expressed in FA-C lymphoblasts that harbored the inactivating FANCC mutation L554P. Wild-type FANCC cDNA completely corrected TNF-α overproduction when expressed in FA-C cells, but none of the mutants we tested were capable of completely normalizing TNF-α production levels (Figure 5B).

Discussion

Most of the scientific literature on FA genes and proteins focuses on the DNA damage and repair response, in which all of the 13 FA proteins play a key role. Some authors have argued that the bone marrow failure phenotype simply devolves from ongoing genotoxicity in the stem cell pool, but recently reports have confirmed that FA proteins participate directly in canonical signaling pathways that influence survival and self-replication of hematopoietic cells. In fact, in the case of FANCC, certain signaling functions of the protein can be formally distinguished from the function of the molecule on protecting...
progenitor cells are affected by in particular by the most well studied of these cytokines, TNF-α, which plays a key role in the pathogenesis of both bone marrow failure and clonal evolution in Fancc−/− mice.9,13 In marrow failure, the TNF-hypersensitive phenotype is likely exacerbated by the overproduction of TNF-α by FA cells,11-13,35 a phenomenon that has been attributed to both transcriptional and posttranslational mechanisms that require activation of MAPK pathways.36 The studies we describe herein clarify a key role of TLR8 in TNF-α overproduction and, in keeping with the studies recently reported by Briot et al,36 we find that p38 MAPK, known to play a role in TLR responses,37-39 is absolutely required for the TNF-α overproduction phenotype (see Figure 4E-G). Both of these studies do confirm that aberrant control of TNF production in FA cells derives at least in part from a transcriptional mechanism,36 and our results with THP-1 blue shFANCC cells indicate that a major mechanism by which p38 influences TNF-α production is by suppressing TNF-α gene transcription. However, TNF-α release can be controlled posttranslationally, and further studies on the relationship of posttranslational control and TLR8 activity is warranted.

In this study a direct comparison of the ubiquitomes of FA-C– and isogenic-complemented lymphoblasts led to the identification of TLR8 uniquely in the ubiquitinated fraction of FA-C cells. In light of the role of TLRs in inducing TNF-α gene expression, we reasoned that the proteomics result might be a report of enhanced TLR8. That is, either TLR8 ubiquitination or the association of TLR8 with an ubiquitinated protein might reflect an enhancement of its activity. We therefore tested the hypothesis that an abnormally high activation state or activation-potential of TLR8 was the mechanism that accounted for the overproduction of TNF-α in FA-C cells. We confirmed this hypothesis.

Specifically, we found that TLR8 (or a TLR8-associated protein) is highly ubiquitinated in mutant FA-C cells and that high-level TNF-α synthesis in mutant cells depended upon TLR8 (Figure 4D) and its downstream signaling intermediates IRAK-1 and IKK-alpha/beta (Figure 4A-B). FANCC-deficient mononuclear phagocytes (THP1 cells) overexpressed TNF-α in response to TLR8 agonists but not other TLR agonists, including the TLR7 agonist imiquimod (supplemental Figure 6),40 and both human FANCC-deficient and murine Fancc-deficient mononuclear phagocytes exhibited hypersensitive TNF-release responses to the TLR8 agonist R848 (Figure 3). Therefore, FANCC functions to antagonize the state of TLR8 activation, and overproduction of TNF-α by FA cells evolves, at least in part, from deregulation of TLR8 activity. We conducted TLR8/TNF complementation studies on FA lymphoblasts by using mutated forms of FANCC (Figure 5) and confirmed that the abnormal TLR8 activation state was not complemented by FANCC mutants that were fully capable of complementing fully in the MMC sensitivity assay.2 This finding confirms that the aberrant activation state of TLR8 in FA-C cells does not simply devolve from ground state genotoxicity.

During the course of these studies we found that although FA lymphoblasts overproduced TNF-α in the ground state, FANCC-deficient mononuclear phagocytes (THP1 blue shFANCC cells) and Fancc-deficient primary splenic macrophages required a TLR agonist (R848) to reveal the TNF-overproduction phenotype and that Fancc-deficient splenic lymphocytes did not exhibit the phenotype. In light of these results and of concerns that EBV transformation of the FA B-cell lines may contribute to the ground state activation of TLR8 in those cells, we focused most of our follow-up studies on mononuclear phagocytes and argue that in light of the murine studies, macrophages represent the relevant cell type vis-à-vis this particular mechanism.

Figure 3. FANCC suppresses TNF production and NF-κB activation induced by TLR8 ligands. Results in THP-1 cells are from 1 of 4 identical experiments. THP-1 Blue cells transduced with control (not targeted) shRNA or with shRNA targeting FANCC were incubated 24 hours with various concentrations of R-848. Expression of the NF-κB reporter was quantified colorimetrically (A), and TNF-α production was quantified by the use of ELISA (B). Dose-response curves shown are representative of 4 (A) and 2 (B) assays. SEAP and TNF-α responses of untransduced cells matched precisely those of cells transduced with the control shRNA (not shown). The specific lentiviral shRNA targeting FANCC (shFANCC) was selected based upon its capacity to induce MMC hypersensitivity and suppress monoubiquitination of FANCC2 in THP1 Blue cells (data are shown in supplemental Figure 5). (C) TNF-α release from primary murine splenic cd11b+ macrophages exposed for 24 hours to multiple doses of R848 showed a 4-fold increase in TNF-α released in cultures of FANCC−/− macrophages. Macrophage-depleted mononuclear cells produced no detectable TNF-α either before or after exposure to R848. Error bars represent mean (± SD).

DNA from cross-linking agent–induced damage.2 The findings described in this study provide an additional example.

Treatment with a variety of hematopoietic suppressive cytokines evinces in FA bone marrow cells a profoundly inhibitory response.2,30,29,34 Survival and replication of hematopoietic stem cells and
Figure 4. Inhibition of the TLR8 pathway suppresses R848 activation of NF-κB in FANCC-deficient cells. THP-1 Blue cells expressing either control (nontargeted shRNA) or FANCC shRNA (target and shRNA sequences are shown in supplemental Table 4) were pretreated for 2 hours with 25μM IKK inhibitor (A) or 25μM IRAK-1/4 inhibitor (B) and then incubated for 24 hours with 30μM R-848. Both inhibitors reduced NF-κB activation in THP-1 Blue cells exposed to R-848. (C) Stable expression of TLR8 shRNA (target and shRNA sequences are shown in supplemental Table 4) in THP-1 Blue cells reduced TLR8 mRNA levels by 73%. Nontargeted shRNA had no effect on TNF-α mRNA. (D) Stable coexpression of TLR8 shRNA with either control (nontarget) or FANCC shRNAs lowered R-848–induced NF-κB activation. Specificity of TLR8 shRNA was confirmed by in experiments that showed that this shRNA did not suppress LPS-induced NF-κB activity (not shown). Error bars represent mean (± SD). (E) NF-κB activation (assessed by quantification of SEAP, y-axis) was quantified in the ground state and in cells treated with R848 in the presence and absence of SB203580, a p38 MAPK inhibitor. The inhibitor suppressed the R848 response in both control and shFANCC cells. (F) TNF-α production was likewise suppressed by SB203580 in THP1 Blue shFANCC cells treated with R848. Control cells not treated with R848 did not produce TNF-α. (G) Likewise, high-level endogenous TNF-α production in FANCC-deficient HSC536 lymphoblasts (control) was also suppressed by SB203580. HSC536/FANCC cells are isogenic FA-C lymphoblasts expressing wild-type FANCC cDNA.
MyD88, Mal, IRAK4, IRAK1, and other TLRs. TLR8 for ubiquitinylation of a TLR8-associated protein. Other potential MMC sensitivity test. 2 (B) HSC536 cells were transduced with wild-type FANCC mutant. All of the alanine mutants are known to fully complement FA-C cells in the MMC sensitivity test. 2 (B) HSC536 cells were transduced with wild-type FANCC cDNA (bar 2), 2 naturally occurring FANCC mutant constructs (bar 1, LS54P and bar 9, M55) and 6 engineered point mutant constructs. With the exception of the partially effective S249A mutant and the completely effective normal cDNA ("wild-type"), which normalized TNF production in FA-C cells, TNF-α levels remain high in all of the isogenic cells expressing the other alanine mutants and M55.

Whether the major increase in TLR8 in antiubiquitin antibody immunoprecipitates reflects excessive ubiquitinylation of TLR8 or a TLR8 binding protein is not yet solved, but there is precedent for direct ubiquitinylation of TLR8 in response to the TLR7/8 agonist 3M-003. However, the design of these previous studies involved overexpression of TLR8 in HEK-293 cells and detection of peptides by mass spectrometry. No ubiquitinylation sites or differential biologic function studies were reported by this group. That there is a discrete increase in the molecular mass of TLR8 in the antiubiquitin antibody immunoprecipitate (Figure 1 and supplemental Figure 1) suggests that TLR8 is ubiquitinated but likely not polyubiquitinated. However, until formal proof of direct ubiquitinylation is developed in further studies, it remains possible, in light of the role played by ubiquitinylation generally in the TLR pathways, 42 that our results can be explained by differential ubiquitinylation of a TLR8-associated protein. Other potential binding partners of TLR8 (or a TLR8-containing complex) include MyD88, Mal, IRAK4, IRAK1, and other TLRs. TLR8 for example, has been shown to interact with TLR7 and TLR9. TLR8, or associated proteins, may also bind to negative regulators, including SIGIRR, ST2L, MyD88s, SOCS1, Tollip, IRAKM, IRAK2/c/d, and TRIAD3A. If an ubiquitinated TLR8-associated protein is present in FA-C cells rather than ubiquitinated TLR8 itself, all of these proteins should be considered as possible targets for this key posttranslational modification. In fact, by using coimmunoprecipitation methods we examined the interaction of 2 FA proteins, FANCA and FANCC, with TLR8 and its signaling intermediates and although we found associations between (1) FANCA and TRAF6 and (2) FANCA and TAK1 (not shown), these interactions were equivalent in FA-C cells and complemented cells. We did not find evidence of FANCC:A20 associations. A20 was of particular interest because it inhibits both TLR and TNF receptor signaling pathways (both of which are perturbed in FA cells) and because its activation affects ubiquitinylation levels of proteins in the TLR and TNF receptor pathways. 36,47 We also found no evidence, when using coimmunoprecipitation, that FANCC or FANCA associated with TLR8, Myd88, IRAK1, IRAK-4, IKK-α, IKK-β, NEMO, IkBα, IkBβ, IκBγ, Triad4A, suppressor of cytokine signaling 1, or IRAK-M.

In summary, although the precise biochemical function of the FANCC protein in the TLR8 pathway has not yet been identified, we have determined that inactivation of FANCC results in an increase in the activation state of TLR8 and a consequent increase in the transcription of TNF-α, a molecule of particular pathophysiologic significance in the bone marrow failure and clonal evolution that characterize FA. Because specific point mutants of FANCC complemented FA-C cells in the MMC assay but did not suppress TNF-α overproduction in FANCC-deficient cells (Figure 5) the TLR8-related abnormality does not simply devolve from ground-state genotoxicity in FA cells. Finally, the TLR8 pathway represents a potentially attractive developmental therapeutic target in FA, and the THP-1 Blue/FANCC shRNA cells provide a convenient tool for high-throughput screening for molecules that suppress this hyperactive pathway in FA-C cells.

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Authorship

Contribution: S.M.V. and G.C.B. designed the project, analyzed all primary data, developed new assays, and wrote the manuscript; S.O. conducted all chromosomal instability testing; J.S. and R.K.R. conducted in vitro assays using murine cells; C.H. conducted gene expression microarrays, analyzed primary gene expression microarray data, and prepared CEL files; J.Y. conducted electromobility shift assay experiments and real-time RT-PCR confirmation of selected microarray results; W.K. conducted all cell-line cultures, including packaging cell lines, conducted MMC sensitivity assays, and conducted retroviral gene transductions; D.C.A. and S.M.V. designed and conducted mass spectrometry experiments and data analysis; N.F.P. coordinated the clinical and laboratory projects in Brazil; D.V.P. prepared and shipped RNA from Brazil to Portland and provided clinical data from Brazilian FA patients; P.A. conducted small molecule screening experiments using THP1 Blue shFANCC cells and analyzed the p38 inhibitor data; and R.P. participated in the design of these studies and accrued study subjects from Brazil.

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References


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TLR8-dependent TNF-α overexpression in Fanconi anemia group C cells

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