A mutation in a CD44 variant of inflammatory cells enhances the mitogenic interaction of FGF with its receptor

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Synovial fluid cells from joints of rheumatoid arthritis (RA) patients express a novel variant of CD44 (designated CD44vRA), encoding an extra trinucleotide (CAG) transcribed from intron sequences flanking a variant exon. The CD44vRA mutant was detected in 23 out of 30 RA patients. CD44-negative Namalwa cells transfected with CD44vRA cDNA or with CD44v3-v10 (CD44vRA wild type) cDNA bound FGF-2 to an equal extent via their associated heparan sulfate chains. However, Namalwa cells, immobilizing FGF-2 via their cell surface CD44vRA, bound substantially more soluble FGF receptor-1 (FGFR-1) than did Namalwa cells immobilizing the same amount of FGF-2 via their cell surface CD44v3-v10. The former cells stimulated the proliferation of BaF-32 cells, bearing FGFR1, more efficiently than did the latter cells. Finally, isolated primary synovial fluid cells from RA patients expressing CD44vRA bound more soluble FGFR-1 to their cell surface-associated FGF-2 than did corresponding synovial cells expressing CD44v3-v10 or synovial cells from osteoarthritis patients. The binding of soluble FGFR-1 to RA synovial cells could be specifically reduced by their preincubation with Ab’s against the v3 exon product of CD44. Hence, FGF-2 attached to the heparan sulfate moiety expressed by the novel CD44vRA variant of RA synovium cells exhibits an augmented ability to stimulate FGFR-1–mediated activities. A similar mechanism may foster the destructive inflammatory cascade not only in RA, but also in other autoimmune diseases.


Introduction

CD44 is a cell surface glycoprotein involved in multiple cellular functions, including cell matrix interactions, cell migration, programmed cell death (apoptosis), or, conversely, cell survival and proliferation. Additionally, CD44 isoforms were shown to exert some of their functions through docking and presentation of cytokines, chemokines, growth factors, and enzymes to their relevant cell surface receptors or substrates (1, 2). Hyaluronic acid is the principal ligand of CD44 (3), but other cell surface or ECM components, such as osteopontin, fibronectin, fibroblast, collagen, and laminin, can interact with this glycoprotein, as well (1). This multifunctionality of CD44 is possible due to the tremendous structural variability of this receptor, derived from its highly complex genetic construction. Theoretically, hundreds of CD44 isoforms can be generated by alternative splicing (4) of ten (mouse) or nine (human) variant exons, designated v1 to v10, inserted in different combinations between the two constant regions consisting of five exons at one end of the molecule and four at the other (1–3). The number of CD44 variants (CD44v) identified so far, however, is limited to a few dozen, detected mostly on epithelial cells, keratinocytes, activated leukocytes, and many types of tumor cells (2). Direct splicing of constant exon 5 to constant exon 16 (thereby skipping all the variant exons) generates the standard CD44 (CD44s), ubiquitously expressed on mesenchymal cells and on all types of hematopoietic cells (1–3). While alternative splicing is a most efficient means of enriching the genetic information stored in a single gene, posttranslational modifications by glycosylation and glycosaminoglycan (GAG) attachments further modify the CD44 protein, allowing greater expansion of its variability and functions (1–3). Indeed, it has been found that heparan sulfate (HS) attached to the v3 exon of v3-containing
CD44 proteoglycans can immobilize HS-binding growth factors, such as hepatocyte growth factor/scatter factor (5–7), VEGF (8), and heparin-binding epidermal growth factor (8–10), as well as FGF-2 (also known as basic FGF) (8, 9, 11), FGF-4 (12), and FGF-8 (12). Previous studies on the mode of action of FGF-2 and its association with HS identified a novel role for cell surface- and ECM-resident HS molecules in the formation of distinct FGF-2-HS complexes that are essential for binding FGF-2 to its cognate receptor (13) and subsequent signal transduction (14, 15). The crucial role of cell surface HS was revealed by the finding that high-affinity receptor binding of FGF-2 is abolished in CHO mutant cell lines defective in the metabolism of GAGs (13). Receptor binding was fully restored upon addition of exogenous heparin (13, 15). Subsequent analysis of the conditions required for high-affinity binding of FGF-2, using soluble recombinant receptors, confirmed the importance of the initial formation of specific FGF-2-HS–receptor complexes for high-affinity binding and activation of FGF receptors (15).

The growth factor–binding function of v3-containing CD44 can support both physiological (e.g., embryonic limb outgrowth) (12) and pathological (e.g., tumor cell motility and growth) (6, 16) activities. We (17–19) and others (20–23) have shown that CD44 targeting by anti-CD44 mAb’s can reduce an experimental tumor growth as well as pathological activities in experimental autoimmune diseases, possibly by disrupting CD44-dependent functions, such as cell migration (24). In most cases, the mAb’s were directed against standard CD44 epitopes, shared by all CD44 isoforms (designated pan-CD44), resulting in targeting of cells engaged in physiological and pathological activities. Therefore, we have focused new efforts on identifying disease-specific CD44 isoforms. Screening for exclusive CD44 variant sequences in synovial fluid cells derived from rheumatoid arthritis (RA) patients, we discovered a novel isoform in which an extra intron-derived trinucleotide is inserted into the CD44v3-v10 variant mRNA of the inflamed joint cells. The expressed RA-associated CD44 variant (designated CD44vRA) bound FGF-2 in a manner promoting high-affinity binding of soluble FGF-receptor-1 as well as the activation of this cell surface receptor.

Methods

Cloning and transfection of human CD44vRA, CD44v3-10, and CD44s. For cloning of human CD44vRA cDNA, the total synovial fluid cell population of RA patients undergoing joint aspiration was isolated. RNA was isolated with the appropriate HRP-conjugated anti-Ig secondary Ab (Jackson Immunoresearch Laboratories, Inc., West Grove, Pennsylvania, USA) for 30 min on ice. The cells were then washed and analyzed with a Flow Cytometer (Becton Dickinson Immunocytometry Systems, San Jose, California, USA).

Western blot analysis. Cells were lysed in NP-40 buffer and 100 µg of protein, run on denaturing SDS-PAGE, and transferred to a PVDF membrane (Millipore Corp., Bedford, Massachusetts, USA). Blots were blocked with 1% BSA in PBS containing 0.1% Tween-20 (PBS-T) and incubated for 1 h with 1 µg/ml polyclonal anti–FGF-2 Ab or 2 µg/ml Hermes-3 anti–pan-CD44 mAb, which was obtained from American Type Culture Collection hybridoma supernatant and purified on a protein-G column. The blots were rewarshed in PBS-T, incubated with the appropriate HRP-conjugated anti-Ig secondary Ab (1:10,000 dilution; Jackson ImmunoResearch Laboratories Inc.) for 45 min, rewarshed in PBS-T, and developed with ECL reagent (Amersham Biosciences Ltd., Little Chalfont, United Kingdom).

Flow cytometry. For flow cytometry, 10⁶ cells were incubated with 3G5 anti-CD44v3 mAb (IgG2b, R&D Systems Inc., Minneapolis, Minnesota, USA), VFF7 anti-CD44v6 mAb (IgG-1; Bender MedSystem GmbH, Vienna, Austria), F-10-44-2 anti–pan-CD44 (known also as anti-CD44s) mAb (IgG-2b; Serotec Ltd., Oxford, United Kingdom), or polyclonal anti–FGF-2 Ab (Serotec Inc.) for 45 min on ice. After extensive washing, the cells were incubated with FITC-conjugated secondary anti-Ig Ab (Jackson Immunoresearch Laboratories Inc., West Grove, Pennsylvania, USA) for 30 min on ice. The cells were then washed and analyzed with a Flow Cytometer (Becton Dickinson Immunocytometry Systems, San Jose, California, USA).

PCR product size was confirmed by agarose gel electrophoresis, sequencing (ABI PRISM 310; PerkinElmer Inc., Wellesley, Massachusetts, USA), and PstI (New England Biolabs Inc., Beverly, Massachusetts, USA) digestion (the nucleotide insertion in CD44vRA introduces a PstI digestion site). The PCR product was excised from the gel, purified, and subcloned into the pGEM vector (Promega Corp.). Positive clones were selected by white/blue screening. Plasmids were purified with a commercial kit (Promega Corp.) subjected to EcoRI/XbaI (New England Biolabs Inc.) double digestion and cloned into the pcDNA3.1 vector (Invitrogen Ltd., Paisley, United Kingdom) in which the gene product was expressed. The plasmid was transfected into the CD44-negative Namalwa Burkitt lymphoma cell line (American Type Culture Collection, Manassas, Virginia, USA) as described (25). For cloning of human CD44v3-10, RNA was isolated from human keratinocytes (a gift from H. Ben-Bassat, Hadassah University Hospital, Jerusalem, Israel); for cloning of human CD44s, RNA was isolated from the HeLa cervical cancer cell line (obtained from American Type Culture Collection), using the above described protocol. Transfection of CD44v3-v10 and CD44s cDNAs as well as of the pcDNA3.1 vector was performed as described above. Accordingly, the transfect- ed Namalwa cells were designated Namalwa-CD44vRA, Namalwa-CD44v3-v10, Namalwa-CD44s, and Namalwa-Neo (Namalwa cells transfected with empty vector).

For cloning of human CD44vRA cDNA, the total synovial fluid cell population of RA patients undergoing joint aspiration was isolated. RNA was isolated with a commercial kit (Promega Corp., Madison, Wisconsin, USA). CD44vRA cDNA was prepared by RT-PCR (PTC-100 Programmable Thermal Controller; MJ Research, Watertown, Massachusetts, USA), using primers representing the constant coding regions of CD44 (see Figure 1): Ex1-sense, 5′-GAATTCGGCCGCCACCATGACGTTTTTGTTGGG-3′; Ex20-antisense, 5′-TCTAGATTACACCGAATCTTCTCATG-3′. PCR product size was
(rhFGF-2) (R&D Systems Inc.) was mixed with 13.44 µg NHS-LC-Biotin (Pierce Chemical Co., Rockford, Illinois, USA) in 5% DMSO and 0.1 M bicarbonate buffer in a total volume of 200 µl. The mixture was incubated for 1 h at 37°C and stored at 4°C. A quantity of 10^6 Namalwa transfectants were incubated with 10 nM biotinylated FGF-2 for 45 min on ice, washed once with or without 0.2 M NaCl and twice with PBS, incubated with streptavidin-phycocerythrin (streptavidin-PE) (Jackson ImmunoResearch Laboratories Inc.) for 30 min on ice, and washed and analyzed by flow cytometry. To block the binding of the FGF-2, cells were incubated with 20 µg/ml heparin or chondroitin sulfate A and C (Sigma Chemical Co., St. Louis, Missouri, USA), together with 10 nM biotinylated FGF-2 for 45 min on ice and washed before FGF-2 binding analysis with streptavidin-PE, as described above. For enzymatic treatment, the cells were incubated with 20 mU/ml heparinase I or with 100 mU/ml chondroitinase ABC (Sigma Chemical Co.) for 2 h at 37°C, washed, incubated with 10 nM biotinylated FGF-2 for 45 min on ice, and rewashed before FGF-2 binding analysis with streptavidin-PE, as described above. To determine the incorporation of FGF-2 into Namalwa transfectants by Western blot analysis, the cells were incubated with 50 nM rhFGF-2 for 45 min on ice, washed with 0.2 M NaCl, and then subjected to SDS-PAGE and blotting with anti–FGF-2 Ab as described (12).

FGF receptor-1 binding assay. A quantity of 5 × 10^6 Namalwa transfectants were washed with PBS and incubated with 0.1, 1, 2, 5, and 10 nM rhFGF-2 for 1 h on ice. The cells were then washed with PBS, mixed with 200 µl conditioned medium containing the soluble FGF receptor-1 (FGFR-1) ectodomain coupled to alkaline phosphatase (FGFR-1-AP) (0.24 OD units/min) (15) for 4 h on ice followed by extensive washing with PBS. AP substrate (pNPP solution; Sigma Chemical Co.) was added to the cells for 3 h at 37°C, and the developed color was analyzed in a spectrophotometer plate reader (Molecular Devices, Sunnyvale, California, USA) at 405 nm. To compare FGFR-1 binding with FGF-2 binding, 10^6 Namalwa transfectants were washed with PBS and incubated with 10 nM biotinylated FGF-2 for 45 min on ice, washed, and reincubated with streptavidin-AP (Jackson ImmunoResearch Laboratories Inc.) for 30 min on ice, and then rewashed extensively with PBS. The pNPP solution was added to the cells for 1 h at 37°C, and the developed color was analyzed in a spectrophotometer plate reader at 405 nm. In addition, 5 × 10^6 synovial fluid cells from the joints of RA or osteoarthritis (OA) patients, as well as primary human keratinocytes, were washed with PBS and incubated in the absence or presence of 10 nM rhFGF-2 for 1 h on ice. Binding of soluble FGFR-1 to these cells was determined as described above. For blocking FGFR-1 binding, 0.01, 0.3, 3, 10 µg/ml anti-CD44v3 mAb, 10 µg/ml anti–pan-CD44 (F-10-44-2), or 10 µg/ml isotype-matched control IgG-2b (Serotec Ltd.) were added to the cells for 1 h on ice. After washing, the binding of FGFR-1 to these cells was determined as described above.

**Figure 1**
Schematic diagram of the CD44 molecule and the trinucleotide CAG insertion in the CD44v3-v10 sequence of RA patients. A schematic of the CD44 genomic map is shown at the top of the figure. The black squares represent the constant exons (designated C1, C2, etc.) at the two ends of the molecule. The white squares represent the variant exons (designated v2, v3, etc.) subjected to alternative splicing. Differential use of the variant exons generates the different CD44 isoforms, e.g., use of exons v3 to v10 in tandem forms the CD44v3-v10. Note that exon v1 is not included in human CD44. The gap between exon v4 and exon v5 indicates the insertion site of CAG, the extra trinucleotide detected in the CD44v3-v10 variant synovial fluid cells (CD44vRA), isolated using RT-PCR from cells removed from the joints of RA patients. Arrows mark the positions and directions of Ex1 sense and Ex20 antisense primers (for sequence, see Methods) used in the above-mentioned RT-PCR. Magnification of the CAG insertion site is shown at the bottom of the figure, indicating the nucleotide sequence at the 3’ end of exon v4 and the 5’ end of exon v5 and its alignment with the published sequence (shown in the boxes) in the same region (29). The drawing in the middle of the figure illustrates the single chain of CD44 proteoglycan, emphasizing the position of the variable region. The HS associated with exon v3 marks the attachment site of a heparan sulfate moiety, the GAG chain that binds to FGF-2 in the CD44 variant.
BaF-32 cell proliferation assay. Namalwa transfectants or synovial fluid cells from the joints of RA patients were fixed with 1% paraformaldehyde (Electron Microscopy Sciences, Washington, Pennsylvania, USA) for 2 h on ice, washed with PBS, and placed on ice for 30 min, rewashed three times, and suspended in RPMI-1640 containing 0.5% FCS. In a modified proliferation assay described (26, 27), the fixed cells were mixed with FGFR-1–transfected BaF-32 cells (a gift of Israel Vlodavsky, Hadassah University Hospital) and 1 nM rhFGF-2. Heparin and rhFGF-2, added to BaF-32 cells as described (8), served as a positive control. BaF-32 proliferation was measured as mitochondrial NADH/NADPH–dependent dehydrogenase activity in a colorimetric assay after 72 hours, using CellTiter 96 AQueous One Solution Reagent (Promega Corp.), and the developed color was analyzed in a spectrophotometer plate reader at 490 nm. To inhibit the proliferation of BaF-32 cells stimulated with Namalwa transfectants, 10 µg/ml of 3G5 anti-CD44v3 mAb or isotype-matched control were added to the cell mixture every 24 hours for 3 days.

Statistical analysis. Data were analyzed using microcomputer programs for one-way ANOVA, followed by the Student t test for unpaired values. P values less than 0.05 were considered significant. The results are expressed as the mean plus or minus SEM. Each experiment was repeated at least three times, all showing similar results.

Results

RA-specific CD44 variant. In a search for disease-specific CD44 isoforms in the synovial fluid cells (comprising approximately 70% polymorphonuclears and macrophages; the remainder are T and B lymphocytes) of RA patients, RT-PCR revealed CD44 variant transcripts, mostly CD44v3-v10, in 44 out of 47 RA patients. The expression of CD44 variants in all synovial fluid cell subpopulations was confirmed by flow cytometry (data not shown). CD44v3-v10 was also identified in normal keratinocytes (28). When the CD44v3-v10 isoform of RA synoviocytes was sequenced, we discovered that it included an extra trinucleotide (CAG) that was illegitimately transcribed from the end of the intron flanking the 5’ end of exon v5, allowing an extra alanine to be encoded, without interfering with the reading frame (Figure 1). Transcripts with an identical sequence change, designated CD44vRA (CD44 variant of RA patients) were detected in 23 out of 30 RA patients whose RT-PCR products were sequenced (not shown). The inclusion of CAG, which introduced a PstI digestion site, was confirmed by PstI digestion. Preliminary evaluation using CD44vRA-specific mAb suggests that the cells expressing the novel variant constitute a major fraction of the RA synovial fluid cells, most likely macrophages and polymorphonuclear cells (data not shown). Aside from the CAG inclusion in the RA CD44 variant, CD44vRA and CD44v3-v10 have an identical sequence, shown by alignment with the entire published CD44 sequence (29), as well as with the CD44 sequence isolated from keratinocytes in our laboratory.

CD44 variants bind FGF-2 to the same extent. Namalwa cells (derived from patients with Burkitt lymphoma), originally lacking CD44, were transfected with (a) standard CD44 cDNA isolated from the HeLa cell line (designated Namalwa-CD44s), (b) CD44v3-v10 cDNA
isolated from primary keratinocytes (Namalwa-CD44v3-v10 or Namalwa-CD44v), (c) CD44v3-v10 cDNA containing the extra CAG, isolated from RA synoviocytes (Namalwa-CD44vRA), or (d) empty vector (Namalwa-Neo). Anti-pan-CD44 mAb (F-10-44-2) stained Namalwa-CD44v4 more intensely than Namalwa-v3-v10 and Namalwa-CD44vRA cells, but did not stain Namalwa-Neo cells. On the other hand, anti-CD44v3 mAb similarly stained Namalwa-CD44v3-v10 and Namalwa-CD44vRA cells, but did not stain Namalwa-CD44s or Namalwa-Neo cells (Figure 2a). These results were confirmed by Western blot analysis with Hermes-3 anti-CD44 mAb (Figure 2b), which showed that extracts of Namalwa-CD44s include only CD44s, whereas extracts of Namalwa-CD44v3-v10, as well as of Namalwa-CD44vRA, contain only the CD44 variant. Namalwa-CD44v3-v10 and Namalwa-CD44vRA cells contain a similar amount of CD44 protein, whereas Namalwa-CD44s cells have a higher level of the ectopically expressed protein.

Heparin-binding growth factors such as FGF-2 can be bound to the V3-associated HS of v3-containing CD44 variants (but not to other variant exons) and then presented autocrinally or paracrinally to the corresponding receptors (30). Flow cytometry revealed that PE-labeled FGF-2 bound at a wide range of concentrations (0.1–20 nM) (not shown) at an extent similar to Namalwa-CD44v3-v10 cells and to Namalwa-CD44vRA cells (Figure 3; 10 nM FGF-2), both in the absence (Figure 3a, inset) and in the presence (Figures 3a, a and b) of 0.2 M NaCl, which reduces the nonspecific binding of FGF-2. In contrast, Namalwa-CD44s cells and Namalwa-Neo cells did not bind FGF-2 at all of the above-indicated concentrations (Figure 3a, a and b; 10 nM FGF-2). The similar binding of FGF-2 to Namalwa-CD44v3-v10 and Namalwa-CD44vRA was confirmed by Western blot analysis with anti-FGF-2 Ab, using cell extracts from Namalwa transfectants expressing the corresponding CD44 variants (Figure 3b). The binding of PE-labeled FGF-2 to Namalwa-CD44vRA (Figure 3c) and to three of its clones (not shown), was blocked by an excess of soluble heparin, and much less so by chondroitin sulfate, indicating that the HS moiety mediates the interaction between FGF-2 and Namalwa-CD44vRA. Similarly, the binding of PE-labeled FGF-2 to Namalwa-CD44vRA (Figure 3d) and to three of its clones (not shown) was inhibited by pretreatment of the cells with heparinase, but not with chondroitinase ABC (Figure 3d), demonstrating that it is the HS moiety of CD44 that mediates the interaction between FGF-2 and Namalwa-CD44vRA.

FGF-2 bound to cell surface CD44vRA interacts with soluble FGFR-1 and enhances FGFR-1–mediated cell proliferation. A soluble FGFR-1 extracellular domain fused to AP showed significantly better binding to Namalwa-CD44vRA cells preincubated with 5 nM FGF-2, than to similarly treated Namalwa-CD44v3-v10 cells (Figure 4a). For comparison, we also assessed the binding of biotinylated FGF-2 to the same transfectants and found that Namalwa-CD44vRA and Namalwa-v3-v10 bound this growth factor to an equal extent (Figure 4a), confirming the results described in Figure 3, a and b. This finding shows that the FGF-2–mediated binding of FGFR-1 to Namalwa-CD44vRA cells is more efficient than that to Namalwa CD44v3-v10 cells, although both transacted cell lines bind similar levels

**Figure 3**

Namalwa-CD44v3-v10 and Namalwa-CD44vRA bind FGF-2 to a similar extent. (a) Flow cytometry. The indicated Namalwa transfectants were incubated with biotinylated FGF-2 in the absence (inset) or presence of 0.2 M NaCl and then analyzed by flow cytometry for their ability to bind this growth factor, detected by staining with streptavidin-PE. Control: Namalwa cells transfected with empty vector (Namalwa-neo) and incubated with biotinylated FGF-2. (b) Western blot analysis. Western blots of cell extracts from Namalwa transfectants with anti-FGF Ab confirmed the flow-cytometry analysis. The Namalwa transfectants were preincubated with FGF-2 before being subjected to cell extraction and gel electrophoresis. The anti-FGF Ab showed that FGF-2 was bound to a similar extent to CD44v3-v10 and CD44vRA, whereas CD44s did not bind FGF-2. Actin, a housekeeping gene product, is equally expressed in all transfectants. (c) Excess soluble heparin blocks the binding of FGF-2 to Namalwa-CD44vRA. Namalwa-CD44vRA cells were coincubated with 5 nM FGF-2, than to similarly treated Namalwa-CD44v3-v10 cells (Figure 4a). For comparison, we also assessed the binding of biotinylated FGF-2 to the same transfectants and found that Namalwa-CD44vRA and Namalwa-v3-v10 bound this growth factor to an equal extent (Figure 4a), confirming the results described in Figure 3, a and b. This finding shows that the FGF-2–mediated binding of FGFR-1 to Namalwa-CD44vRA cells is more efficient than that to Namalwa CD44v3-v10 cells, although both transacted cell lines bind similar levels...
of FGF-2. Repeating the direct binding experiment, this time using several different concentrations of FGF-2, gave us an opportunity to learn also about the saturation binding kinetics of this reaction (Figure 4b). From the initial rates of binding under conditions where the HP-ligand binding sites for FGF-2 are saturated and show a typically high affinity and specific binding profile, we could estimate an apparent dissociation constant of about 100 pM for both CD44vRA and CD44v3-v10-associated FGF-2. This is well within the known range for FGF-2 binding to cells expressing FGFR-1. The maximum binding capacity, however, as determined by the values of FGFR-1 bound at saturating ligand concentrations, was about twofold higher for the CD44vRA isoform. Overall, the kinetic study confirms the results presented in Figure 4a and points to the molecular mechanism whereby the vRA mutation induces a specific increase in HS-mediated FGFR-1 binding and activation.

To analyze the mitogenic activity of cell surface-bound FGF-2, we assessed the ability of fixed Namalwa transfectants incubated with FGF-2 (termed FGF-2–immobilized fixed Namalwa transfectants) to stimulate the proliferation of BaF-32 cells.
expressing FGFR-1 (8). FGF-2–immobilized fixed Namalwa-CD44vRA cells stimulated a proliferative response in cocultured BaF-32 cells that was similar to (Figure 4c, inset), or as shown in different experiments, lower than (Figure 4, c and d) that of the BaF-32 cells incubated with FGF-2 and heparin (positive control). In all three experiments, however, FGF-2–immobilized fixed Namalwa-CD44vRA cells stimulated BaF-32 cells to a greater extent than did the corresponding Namalwa-CD44v3-v10 cells. We further show (Figure 4d) that anti-CD44v3 mAb reduced the proliferative response of BaF-32 cells stimulated with FGF-2–immobilized fixed Namalwa-CD44vRA cells (bars 6, 7, and 8) or with the corresponding Namalwa-CD44v3-v10 cells (bars 3, 4, and 5). BaF-32 cells incubated with FGF-2 alone or heparin alone showed only a background level of cell proliferation, similar to the proliferation rate of BaF-32 cells stimulated with FGF-2–immobilized fixed Namalwa-CD44s or Namalwa-Neo cells (Figure 4c and inset). We also demonstrate (Figure 4d; bars 13, 14, and 15) that fixed RA synovial fluid cells, endogenously loaded with FGF-2 (see Figure 5b), can also stimulate the proliferation of BaF-32 cells. This proliferation was reduced following the addition of anti-CD44v3 mAb (not shown).

Enhanced binding of FGFR-1 to cells isolated from synovial fluid of RA patients. Based on the enhanced binding of FGFR-1 to Namalwa cells expressing CD44vRA, it was important to test and confirm these findings in the authentic cells isolated from the synovial fluid of patients with joint diseases. Flow-cytometry analysis of 18 RA patients and 6 OA patients showed that although joint synovial fluid cells of RA and OA patients expressed pan-CD44 proteoglycan to a nearly equal extent, only RA synovial fluid cells expressed CD44v3 and CD44v6 epitopes (a representative data in Figure 5a). Furthermore, immunostaining with anti–FGF-2 Ab to detect the presence of FGF-2 on the cell surface of synovial fluid cells revealed that RA and OA cells contain similar levels of FGF-2 (Figure 5b, insets). RA synovial fluid cells, however, bound soluble FGFR-1 more

Figure 5
Synovial fluid cells from RA patients bind soluble FGFR-1. (a) CD44 expression on synovial fluid cells from RA and OA patients. Cells collected from the joints of an RA or an OA patient were analyzed by flow cytometry using anti–pan CD44 mAb (F-10-44-2), anti-CD44v3 mAb, or anti-CD44v6 mAb. The first histogram in each panel shows staining with second Ab alone. Similar flow-cytometric histograms were recorded in 11 samples of RA patients and 6 of OA patients. (b) Enhanced binding of soluble FGFR-1 to synovial fluid cells of RA patients. Cells from the joints of 11 RA and 6 OA patients were incubated with soluble FGFR-1 conjugated to AP, in the presence (not shown) or absence (b) of FGF-2. The interaction of the FGFR-1 with the FGF-2, bound to the joint cells, was detected using an AP substrate at OD 405. Similar results were observed in the presence of FGF-2 (not shown). Insets: Binding of anti–FGF-2 Ab to joint cells of RA and OA patients to assess the endogenous FGF-2 inclusion in these cells. The first histogram in each inset shows staining with second Ab only (goat anti-rabbit Fab’-FITC). Equal levels of endogenous FGF-2 in RA and OA cells were observed in samples of 11 RA and 6 OA patients. (c) Enhanced binding of soluble FGFR-1 is dependent on CD44vRA expression on RA synovial fluid cells. The binding of FGFR-1 to three patient samples (RA12, RA13, and RA14) of RA synovial fluid cells expressing CD44vRA, one patient sample (RA15) of RA synovial fluid cells expressing CD44v3-v10, and two samples of primary human keratinocytes (Ker-1 and Ker-2) expressing CD44v3-v10 were analyzed as indicated in b. Note that the keratinocytes were loaded with FGF-2 because, unlike synovial fluid cells, they do not contain the endogenous growth factor. Statistical analysis of the principal groups is shown.
dose-dependent manner by anti-CD44v3 mAb (Figure 6a) but not by F-10-44-2 anti-pan CD44 mAb (Figure 6b), indicating that FGFR-1 binding is targeted to the v3 exon of cell surface CD44.

**Discussion**

A novel CD44 variant CD44vRA was found in the synovial fluid cells from 23 out of 30 examined RA patients. When expressed in a model of Namalwa cells or in original synovial fluid cells of RA patients, it conferred an enhanced capacity to bind soluble FGFR-1 via FGF-2 immobilized on the cell surface CD44 isoform and to stimulate the proliferation of BaF-32 cells bearing FGFR-1. Because FGF-2 was shown to play an active role in the inflammatory response and in disease-associated angiogenesis (see below), we suggest that enhanced binding and activation of FGFR-1 by CD44vRA-expressing synovial fluid cells of RA patients play a role in the RA pathology.

The illegitimate transcription of the intronic CAG flanking exon v5 of CD44vRA is presumably a consequence of misregulation of the splicing machinery in this molecule. Unknown genetic or environmental factors or a combination of both may modify the relative abundance, tissue distribution, or activity of serine-arginine, or of heterogeneous nuclear ribonucleoprotein splicing factors that antagonistically control the differential splicing (31). The CAG-containing splicing junction may be particularly susceptible to such changes, resulting in the CAG inclusion in the CD44 mRNA sequence. A similar mechanism may influence the CD44 splicing machinery of other autoimmune diseases. In this context, it should be mentioned that mutations located in noncoding regions such as those affecting 5’ and 3’ splice sites, branch sites, or polyadenylation signals, are frequently (∼15%) the cause of genetic diseases (32).

For efficient autocrine or paracrine presentation to the relevant neighboring receptors, FGF-2, like other heparin-binding growth factors, must be assembled on cell surface HS proteoglycans (13, 33). The proteoglycan nature of CD44 of cell surfaces is well established (8, 9, 30). CD44 includes seven potential consensus single serine-glycine (SG) or double SGSG assembly sites for glycosaminoglycans (GAGs), such as HS or chondroitin sulfate. It was found, however, that the assembly of HS is determined by eight amino acids located downstream from the exon v3 SGSG motif. Because exon v3 of the CD44 proteoglycan is the only one containing this sequence in the context of the SG or SGSG motif, HS assembly is confined to this exon (30). The sequestering of HS-binding growth factors on the v3 exon of CD44 proteoglycan has been well documented (5–12). Moreover, Sherman and colleagues (12) demonstrated that the addition of anti-CD44v3 mAb to limb bud mesenchymal cells cocultured in the presence of FGF-8, with UV-irradiated Namalwa cells expressing CD44v3-v10, markedly reduced the proliferation of the FGFR-bearing mesenchymal cells. Similarly, we show here that

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**Figure 6**

Synovial fluid cells from RA patients bind soluble FGFR-1 in a CD44v3-dependent manner. (a) Binding of soluble FGFR-1 to synovial fluid cells of RA patients is CD44v3 associated. Synovial fluid cells from three RA patients (RA6, RA8, RA11) were incubated with soluble FGFR-1 conjugated to AP in the presence of medium (bar 1) isotype-matched control immunoglobulin (bar 2), or 1 µg (bar 3), 300 ng (bar 4), 30 ng (bar 5), and 1 ng (bar 6) anti-CD44v3 mAb. The interaction of FGFR-1 with FGF-2, bound to the joint cells, was analyzed as indicated in Figure 5b. Anti-CD44v3 mAb reduced the binding of soluble FGFR-1 to the synovial fluid cells in a dose-dependent manner. The highest concentration (1 µg) of anti-CD44v3 mAb reduced the binding of soluble FGFR-1 to synovial fluid cells in a dose-dependent manner. (b) Anti-CD44v3 mAb, but not anti–pan CD44 mAb (F-10-44-2), directed against a constant epitope inhibits the binding of soluble FGFR-1 to synovial fluid cells of RA patients. Synovial fluid cells from three RA patients (RA2, RA3, RA6) were incubated with soluble FGFR-1 conjugated to AP in the presence of medium (bar 1), isotype-matched control immunoglobulin (bar 2), 1 µg anti-CD44v3 mAb (bar 3), and 1 µg anti–pan CD44 mAb (bar 4). The interaction of the FGFR-1 with FGF-2 bound to the joint cells, was analyzed as described in Figure 5b.

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intensively than did OA synovial fluid cells, regardless whether FGF-2 was added to the cells (not shown) or not (Figure 5b). Addition of heparin at saturated concentrations (2 and 5 µg/ml) to OA cells did not enhance the binding of FGFR-1 to these cells (not shown). Moreover, RA synovial fluid cells expressing the CD44vRA variant (detected by PstI digestion) bound soluble FGFR-1, as indicated by analyzing CD44vRA-positive samples from three RA patients. In contrast, RA synovial fluid cells and FGF-2 externally loaded primary keratinocytes expressing the wild-type CD44v3-v10 (our finding and ref. 28), bound considerably less of this receptor, as indicated by analyzing CD44v3-v10-positive samples from one RA patient and two keratinocyte donors (Figure 5c). The binding of soluble FGFR-1 to synovial fluid cells of RA patients was inhibited in a
anti-CD44v3 mAb specifically reduces the binding of soluble FGFR-1 to synovial fluid cells expressing the RA variant of CD44 (Figure 6) and the proliferation of BaF-32 cells after stimulation with Namalwa CD44v transfectants (Figure 4d) or RA synoviocytes (not shown). Cumulatively, these findings unequivocally show that the HS-binding growth factor is attached to the v3 exon. HS attached to the v3 of CD44 proteoglycan, which can potentiate the binding of growth factors, was detected on, among other cells, inflamed synovial membrane macrophages, and such expression is greatest in cells immobilizing high levels of FGF-2 (8).

The trinucleotide (CAG) insertion between exon v4 and exon v5, interpolating an alanine residue at position 346, constitutes the only difference between CD44vRA and CD44v3-v10 transcripts (Figure 1). Interestingly, inclusion of this single amino acid is sufficient to confer upon FGF-immobilized CD44vRA the capacity for enhanced binding of soluble FGFR-1 and augmented mitogenic activity, without any significant change in the overall FGF-2 binding ability of CD44-associated HS chains. These findings imply that, although quantitatively CD44vRA and CD44v3-v10 display a similar HS-dependent FGF-2 binding capacity, qualitatively, FGF-2-immobilized CD44vRA more effectively promotes interaction with FGFR-1. Two alternative mechanisms, not necessarily mutually exclusive, may account for this observation. It is conceivable that the local structural change induced by alanine insertion may indirectly affect the structure of the more distally v3 exon-attached HS chain so that it can now act as a superior growth factor activator. This, in turn, could result from a local effect on the attachment or recruitment of HS-modifying enzymes responsible primarily for the HS sulfation pattern, including charge density and distribution, critical to its activating or inhibitory functions. Both activities have been implicated in the control of growth factor function in vitro (34) and in vivo (35).

Alternatively, or in addition, the change in the primary sequence of the CD44vRA protein could modify the orientation of existing protein-associated HS structures, enabling them to bind FGF-2 and its receptor in a superior functional form. Analysis of the binding kinetics strongly implies the presence of additional effective and specific binding sites for FGFR-1 on CD44vRA, compared with CD44v3-v10, rather than a change in affinity, as responsible for the new steady-state level obtained. The availability of additional binding sites for FGFR-1 on CD44vRA-associated HS could result from exposure of an existing site that is masked in the CD44v3-v10 isoform by local conformational changes in the core protein or the creation of new binding sites with similar affinity, specific to the CD44vRA-associated HS. The conformational change related to the modified sequence of CD44vRA is demonstrated, using a mAb that was found to bind at higher affinity to Namalwa cells expressing CD44vRA than to Namalwa cells expressing the wild-type CD44v3-v10, although the two transfectants expressed the same level of CD44 (our unpublished observations).

FGF-2, generated at least partly by the joint synoviocytes of RA patients (36), is tightly associated with the exacerbation of RA pathology. Elevated levels of FGF-2 were detected in the synovial tissue (36), synovial fluid (37), and serum (38) of RA patients. FGFR 1, a high-affinity receptor for FGF-2, is also expressed in a variety of RA synovial cells, including endothelial cells (39) (reviewed in ref. 40). The interaction of this growth factor with its cell surface receptor induces, directly or indirectly, osteoclastogenesis and osteocartilaginous destruction in the joints of RA patients (41), as well as in those of rats with adjuvant-induced arthritis (AIA) (37). It was further shown (37) that in rats with AIA, FGF-2 induces, either directly or via stimulation of VEGF release (42), the formation of a new vascular network that supports the inflammatory process.

Although OA synovial fluid cells lack the v3 exon product, they immobilize FGF-2 to nearly the same extent as RA cells (Figure 5b, insets). This finding suggests that OA synovial fluid cells bind FGF-2 via HS of non-CD44 proteoglycan origin. Nevertheless, FGF-2 bound on OA cells is less potent than this growth factor bound on RA cells, because only RA cells show enhanced binding of FGFR-1 (Figure 5b). We can assume, therefore, that binding of FGF-2 to the v3-attached HS of CD44 proteoglycan (expressed on RA, but not on OA cells) markedly improves the potency of this growth factor.

In conclusion, we have identified, we believe for the first time, a novel variant of CD44 in the synovial fluid cells of RA patients. This exclusive CD44vRA variant interacts with FGF-2 via v3-attached HS in a way that allows enhanced binding and activation of FGFR-1, an event that may contribute to the RA inflammatory process. The exclusive structure of synoviocyte CD44vRA should also allow selective targeting, e.g., with mAb, that could interfere with the vicious inflammatory cascade, leading to joint destruction.

Acknowledgments

This work was supported by the Society of Research Associates of the Lautenberg Center (New York, New York, USA) and by a grant from ProChon Biotech Ltd. We thank Israel Vlodavsky for the BaF-32 cells, Alexandra Mahler for her editorial assistance, and Sharon Saunders for typing the manuscript.
