Roles of USF, Ikaros and Sp proteins in the transcriptional regulation of the human reduced folate carrier B promoter

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The hRFC (human reduced folate carrier) is ubiquitously but differentially expressed in human tissues and its levels are regulated by up to seven non-coding regions (A1, A2, A, B, C, D and E) and at least four promoters. For the hRFC-B basal promoter, regulation involves binding of Sp (specificity protein) transcription factors to a critical GC-box. By transiently transfecting HT1080 cells with 5′- and 3′-deletion constructs spanning 1057 bp of upstream sequence, a transcriptionally important region was localized to 158 bp flanking the transcriptional start sites. By gel shift and chromatin immunoprecipitation assays, USF (upstream stimulatory factor) promoter and Sp1 and Ikaros-related proteins were bound to consensus elements (one E-box, two GC-box and three Ikaros) within this region. The functional importance of these elements was confirmed by transient transfections of HT1080 cells with hRFC-B reporter constructs in which they were mutated, and by co-transfections of Drosophila Mel-2 cells with wild-type hRFC-B promoter and expression constructs for USF1, USF2a, Sp1 and Ikaros 2 and 8. Both USF1 and Sp1 proteins transactivated the hRFC-B promoter. Sp1 combined with USF1 resulted in a synergistic transactivation. Identical results were obtained with USF2a. Ikaros 2 was a repressor of hRFC-B promoter activity whose effects were partly reversed by the dominant-negative Ikaros 8. In HT1080 cells, transfection with Ikaros 2 decreased endogenous hRFC-B transcripts, whereas USF1 and Sp1 increased transcript levels. Ikaros 2 also decreased reporter gene activity and levels of acetylated chromatin associated with the endogenous promoter. Collectively, these results identify transcriptionally important regions in the hRFC-B promoter that include multiple GC-box, Ikaros and E-box elements. Our results also suggest that cooperative interactions between transcription factors Sp1 and USF are essential for high-level hRFC-B transactivation and imply that these effects are modulated by the family of Ikaros proteins and by histone acetylation.

Key words: chromatin, Ikaros, methotrexate, reduced folate carrier, Sp1, upstream stimulatory factor (USF).

INTRODUCTION

Reduced folates are essential cofactors for one-carbon transfer reactions in anabolic pathways leading to the synthesis of purines, thymidylate, serine and methionine [1]. Since mammals lack the ability to synthesize reduced folates de novo, cellular uptake of these derivatives is essential for cell growth and survival. The primary route for the membrane transport of reduced folate cofactors is via the RFC (reduced folate carrier) [2,3].

RFC is also responsible for the cellular uptake of ‘classical’ antifolate cancer chemotherapeutic drugs, e.g. Mtx (Methotrexate), Tomudex and Pemetrexed [2,4]. Impaired transport frequently occurs accompanying exposure of human and rodent cells to antifolate drugs in vitro [5–7] and in murine tumour cells in vivo after chemotherapy with Mtx [8]. Furthermore, alterations in the levels of hRFC (human RFC) were associated with Mtx resistance in leukaemias [9] and solid tumours [10] from patients. In B-precursor ALL (acute lymphoblastic leukaemia) lymphoblasts, hRFC expression spans a wide range and is proportional to Mtx uptake [11].

Previous studies suggested a complex regulation of hRFC gene expression [12–16]. We reported recently that hRFC is ubiquitously but differentially expressed in human tissues and cell lines, and identified up to seven non-coding regions (designated A1, A2, A, B, C, D and E) for the hRFC gene in assorted human tissues spanning approx. 35 kb upstream of the major translational start [15]. After splicing, at least 18 potential hRFC transcripts would be generated with unique 5′-UTRs (5′-untranslated regions) fused to the same open reading frame. By analogy with other multi-promoter genes, the unique non-coding exons for hRFC are probably transcribed from distinct promoters. To date, promoter activity has been confirmed for the 5′ regions proximal to the non-coding A2, A, B and C regions [13–16].

To better understand the basis for varied levels of hRFC in human tissues and tumours, we have begun to identify and characterize critical transcription factors involved in regulating the hRFC promoters. We have reported previously that the hRFC-A and -B basal promoters are regulated by different families of transcription factors, including the bZip superfamily (e.g. c-Jun/c-Fos and Creb1/ATF1) and the Sp (specificity protein) family of DNA-binding proteins (e.g. Sp1 and Sp3) respectively [14]. Our results imply that cell-specific expression of these transcription factors can profoundly influence patterns of promoter and 5′-UTR usage at the basal promoter level. However, the net effect of these factors could be overshadowed or enhanced by transcription factor binding to other regulatory elements both upstream and downstream of the minimal hRFC promoter regions, as well as by the state of the surrounding chromatin structure. For instance, promoter A is transactivated by AP2 and Sp1 in co-transfection experiments, via binding to upstream elements [17]. The effects

Abbreviations used: ALL, acute lymphoblastic leukaemia; BST2, bone marrow stromal cell antigen 2; ChIP, chromatin immunoprecipitation; HDAC, histone deacetylase; Mtx, methotrexate; RFC, reduced folate carrier; hRFC, human RFC; 5′-RACE, 5′-rapid amplification of cDNA ends; Sp1, specificity protein 1; USF, upstream stimulatory factor; 5′-UTR, 5′-untranslated region; wt, wild-type.

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of Sp1 on promoter A activity were different, depending on the identity of the bZip protein bound to the basal region.

In the present study, we describe extensive studies of promoter B, a major hRFC promoter in immortalized cell lines and primary leukaemias from patients [15,16]. Our results document transcriptionally important regions in the hRFC-B promoter that include GC-box, E-box and Ikaros elements. Our results also suggest that co-operative interactions between the transcription factors Sp1 and USF (upstream stimulatory factor) are essential for high-level hRFC-B transactivation and imply that these effects can be modulated by the family of Ikaros proteins and by histone acetylation/deacetylation.

**MATERIALS AND METHODS**

**Chemicals and reagents**

[γ-32P]ATP (3000 Ci/mmol) was purchased from PerkinElmer (Boston, MA, U.S.A.), synthetic oligonucleotides were from Sigma-GenoSys Biotechnologies (The Woodlands, TX, U.S.A.), Lipofectin® was from Gibco Life Technologies (Gaithersburg, MD, U.S.A.), and restriction and modifying enzymes, reporter gene vectors (pGL3-Basic and pRLSV40) and other molecular biologicals were obtained from Promega (Madison, WI, U.S.A.). The pCMV-Sp1, pPacO and pPacSp1 plasmid constructs were from hRFC-B/luciferase reporter constructs gifts from Dr Robert Tjian (University of California, Berkeley, CA, U.S.A.). The pPacSp3 construct was provided by Dr Guntram Suske (Philippus-Universitat, Marburg, Germany).

**Cell culture**

The HT1080 human fibrosarcoma cell line was obtained from the A.T.C.C. (Rockville, MD, U.S.A.). HT1080 cells were grown in RPMI 1640 medium with 10% (v/v) heat-inactivated iron-supplemented calf serum (Hyclone, Logan, UT, U.S.A.), 2 mM L-glutamine, 100 units/ml penicillin and 100 µg/ml streptomycin, in an atmosphere of 5% CO2 and 95% air at 37 °C. Drosophila Mel-2 cells were purchased from Invitrogen and were maintained at ambient temperature in Schneider’s insect medium, supplemented with 10% (v/v) fetal bovine serum (Gibco Life Technologies) and 2 mM glutamine plus antibiotics.

**hRFC-luciferase reporter constructs**

The full-length hRFC-B/− 5422 construct [positions −5422 to −4364; the numbering is relative to the ATG translation start in exon 1] and is based on the nucleotide sequence of chromosome 21 contig HS21C102 (accession number AL163302) was described by Whetstone and Matheyer [14] (previously designated as hRFC-B/− 2016). 5′ deleted hRFC-B constructs were prepared from hRFC-B/− 5422 by successive exonuclease III and S1 nuclease digestions (hRFC-B/− 4493 and hRFC-B/− 4448) and restriction endonuclease digestions (hRFC-B/− 4693(Apal), hRFC-B/− 4615(BsrEII), hRFC-B/− 4590(BsmBI) and hRFC-B/− 4467(BssHII)). A 3′-deletion was generated in hRFC-B/− 5422 by restriction endonuclease digestion with BssHII. All deletion constructs were verified by automated DNA sequencing at the Wayne State University Center of Molecular Genetics Sequencing Facility.

Specific DNA-binding sites in promoter B were mutated by PCR, using the hRFC-B/− 5422 construct as a template. All of the mutant constructs [hRFC-B/− 4501MIK(a), hRFC-B/− 4565 MIK(b), hRFC-B/− 4573MIK(c), hRFC-B/− 4534MIK(b) and hRFC-B/− 4434ME(a)] were generated by splice-overlap-extension (‘Soeing’) PCR [18] using the mutant primers shown in Table 1. Primary PCR amplifications used the mutant antisense primers with the upstream ProB (Xhol) primer [5′-CGGGATCCGTACAGGTTACGACCGGCTCCTCC-TCCGATTTCGTGAGCAG-3′] and the mutant sense primer with the downstream 33–54 wt primer [5′-CTGCCTCTGAGGTTACGACCGGCTCCTCC-TCCGATTTCGTGAGCAG-3′]. The primers were mixed and used for the secondary PCR amplifications with the ProB (Xhol) and 33–54 wt (wild-type) primers. The following PCR conditions were used: (i) for the primary PCR amplification, 95 °C for 15 s, 65 °C for 45 s and 72 °C for 40 cycles; and (ii) for the secondary PCR amplification, 95 °C for 15 s, 68 °C for 45 s and 72 °C for 40 cycles. For all mutants, the secondary PCR products were isolated from 2% (v/v) agarose and digested with Xhol and HindIII. The digested fragments were subcloned into pGL3-Basic in the sense orientation and each mutation was verified by automated DNA sequencing.

**Preparation of transcription factor expression constructs**

The pCDNA3-USF1, pPacUSF1 and pPacUSF2a constructs were prepared as described by Ge et al. [20]. For preparing the Ikaros 2 and 8 expression constructs, Ikaros 2 and 8 coding sequences were PCR-amplified from cDNA prepared from CCRF-CEM cells using forward (5′-GCACCACGCTTATGGTTGCTGGAAACC-3′) and reverse (5′-GGGTAAGCGGTGGCTCGCCGCGAC-3′) primers. The amplicons were isolated and mixed for the secondary PCR amplifications with the ProB (Xhol) and 33–54 wt (wild-type) primers. The following PCR conditions were used: (i) for the primary PCR amplification, 95 °C for 15 s, 65 °C for 45 s and 72 °C for 40 cycles; and (ii) for the secondary PCR amplification, 95 °C for 15 s, 68 °C for 45 s and 72 °C for 60 s for 40 cycles. For all mutants, the secondary PCR products were isolated from 2% (v/v) agarose and digested with Xhol and HindIII. The digested fragments were subcloned into pGL3-Basic in the sense orientation and each mutation was verified by automated DNA sequencing.

**Transient transfections and reporter gene assays**

HT1080 cells were transiently transfected with 3 μg of hRFC-B promoter constructs (pGL3-Basic) and 25 ng of pRLSV40 plasmid using Lipofectin® [14]. After 48 h, lysates were prepared, and firefly luciferase activities were assayed using the dual

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Table 1 Primers used to create mutant promoter B constructs

<table>
<thead>
<tr>
<th>Construct</th>
<th>Position</th>
<th>Antisense primer*</th>
<th>Sense primer*</th>
</tr>
</thead>
<tbody>
<tr>
<td>hRFC-B/− 4573MIK(c)</td>
<td>−4573 to −4554</td>
<td>5′-CCCADGGGCAACCCCGAC-3′</td>
<td>5′-GTGGGGGTGGCGTCGACG-3′</td>
</tr>
<tr>
<td>hRFC-B/− 4565MIK(b)</td>
<td>−4565 to −4545</td>
<td>5′-CCCADGGGCAACCCCGAC-3′</td>
<td>5′-GTGGGGGTGGCGTCGACG-3′</td>
</tr>
<tr>
<td>hRFC-B/− 4534MIK(b)</td>
<td>−4534 to −4506</td>
<td>5′-GGGTAAGCGGTGGCTCGCCGCGAC-3′</td>
<td>5′-GTGGGGGTGGCGTCGACG-3′</td>
</tr>
<tr>
<td>hRFC-B/− 4501MIK(a)</td>
<td>−4501 to −4475</td>
<td>5′-GGGTAAGCGGTGGCTCGCCGCGAC-3′</td>
<td>5′-GTGGGGGTGGCGTCGACG-3′</td>
</tr>
<tr>
<td>hRFC-B/− 4434MAE(a)</td>
<td>−4434 to −4414</td>
<td>5′-CCCACGGGCATCGACCTGCCGGA-3′</td>
<td>5′-CTCCAGGAGTCTCAGGCCGAC-3′</td>
</tr>
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* Mutant antisense and sense primers were used for ‘Soeing’ PCR [18] to construct the mutant promoter B constructs, as described in the Materials and methods section. The mutated nucleotides are underlined.
Luciferase kit (Promega) in a T20/20 luminometer. Firefly luciferase activity was normalized to Renilla luciferase activity.

Fugene™6 reagent (Roche, Indianapolis, IN, U.S.A.) was used for transfections into Drosophila Mel-2 cells. For co-transfections, 1 µg of the hRFC-B/−5422 construct in pGL3-Basic was combined with the following expression constructs: 25–100 ng of Sp1 or Sp3 (pPacSp1 or pPacSp3 respectively); 125–500 ng of USF1 or USF2a (pPacUSF1 and pPacUSF2a respectively); and/or 100 or 200 ng of the Ikarsos 2 and 8 isoforms (pPacIk2 and pPacIk8 respectively). Cells were harvested after 24 h for luciferase assays using the Single Luciferase Assay System (Promega). Luciferase activities were normalized to cellular protein contents.

For all transfections, three or more experiments were performed in duplicate.

**Gel mobility-shift assays**

Nuclear extracts from HT1080 cells were prepared as described previously [14], and 12–15 µg of nuclear extract were used in each binding reaction. Gel shift assays were performed exactly as described previously [14]. The [γ-32P]ATP end-labelled probes and competitive mutant oligonucleotides used in the binding reactions are summarized in Table 2. Competition experiments included 150-fold molar excess of the unlabelled wt hRFC oligonucleotides, hRFC oligonucleotides containing mutations in the DNA-binding sites for specific transcription factors or oligonucleotides containing consensus transcription elements, including IKBS1 and IKBS4 [19], GC-box and E-box (both GC-rich reagent and polymerase (Roche). An unrelated gene [BST2 (bone marrow stromal cell antigen 2)] was also amplified with forward (5'-CCGGACCCGAGCGAGCCGTCGCTGATT-3') and reverse (5'-CGGAGGGAGGAGGGCCACGACAGTCTGCA-3') primers and 18 S RNA levels were calculated from their respective standard curves; hRFC-B and 18 S RNA amplicons, which were amplified with the above primers, cloned into pGEM T-Easy vector and linearized with Sall and Apol respectively. Levels of hRFC-B transcripts and 18 S RNA were calculated from their respective standard curves; hRFC-B levels were normalized to 18 S RNA. Real-time PCR results were expressed as relative transcript levels ± S.E.M. for three experiments.

**ChIP (chromatin immunoprecipitation) assay**

ChIP assays were performed in HT1080 cells as described previously [20,21] with antibodies to Sp1 (Active Motif), USF1 (Santa Cruz Biotechnology), Ikars (sc9861; Santa Cruz Biotechnology) and acetyl histone H3 (H06599; Upstate Biotechnology, Lake Placid, NY, U.S.A.). Standard PCR for the hRFC-B promoter region was performed with forward (5'-CCGGACCCGAGCGAGCCGTCGCTGATT-3') and reverse (5'-CGGAGGGAGGAGGGCCACGACAGTCTGCA-3') primers spanning positions −4560 to −4353 using a GC-rich reagent and polymerase (Roche). An unrelated gene [BST2 (bone marrow stromal cell antigen 2)] was also amplified with forward (5'-CCGGACCCGAGCGAGCCGTCGCTGATT-3') and reverse (5'-CGGAGGGAGGAGGGCCACGACAGTCTGCA-3') primers to validate the specificity of the ChIP assays. PCR conditions were designed to ensure linearity and were 95 °C for 5 min, followed by 32 cycles of 95 °C for 30 s, 62 °C for 45 s and 72 °C for 45 s.

**RESULTS**

**Deletion analysis of the hRFC-B promoter**

The hRFC-B promoter is a major promoter for hRFC in immortalized cells and primary leukaemia cells [15,16]. We previously localized the minimal promoter region for hRFC-B to within 46 bp (positions −4493 to −4448) and demonstrated an important regulatory role for the Sp family of transcription factors in binding to a GC-box consensus element [designated GC-box (a)] within this region [14]. Since Sp1/Sp3 proteins are ubiquitously expressed [22], significant differences in hRFC-B transcripts between tissues [15] probably reflect contributions of other cis regulatory elements and transcription factors in regulating promoter activity.
In vitro and in vivo binding of USF, Sp1 and Ikaros transcription factor families to the hRFC-B promoter

The regions that showed major fluctuations in promoter activity with progressive 5' and 3' deletions (Figure 1A) were obvious candidates for in vitro analyses of transcription factor binding. Sp1/Sp3 binding to GC-box (a) was previously confirmed on gel shifts using an oligonucleotide probe from positions −4493 to −4448 [14], not including the Ikaros (a) binding site. Additional gel shifts were performed with nuclear extracts prepared from HT1080 cells and double-stranded oligonucleotide probes spanning positions −4573 to −4414 (Figures 2A–2E) and including the putative transcription factor binding sites shown in Figure 1(B). A total of eight DNA–protein complexes (A–H) were identified that were effectively competed by unlabelled hRFC oligonucleotides, establishing binding specificity.

With the hRFC-B/−4573 (positions −4573 to −4554), hRFC-B/−4565 (positions −4565 to −4545) and hRFC-B/−4501 (positions −4501 to −4475) oligonucleotides, a series of complexes (designated A, B, C, F and G in Figure 2, lanes 2, 7, and 18) were identified that were effectively competed with IBKS1 (not shown) and IBKS4 Ikaros consensus oligonucleotides (lanes 4, 9 and 20). For complexes A, B and C, but not F or G, competition was at least partly abolished with hRFC oligonucleotides including mutant Ikaros binding sites [hRFC-B/−4573MIK(c), hRFC-B/−4565MIK(b) and hRFC-B/−4501MIK(a); mutant sequences shown in Table 2] (lanes 5, 10 and 22). These results strongly imply that complexes A, B, C, F and G all involve binding by DNA–binding members of the Ikaros family of proteins (Ikaros 1, 2 and 3) [19] to the Ikaros (c), (b) or (a) sequence elements depicted in Figure 1(B). The failure of the mutant hRFC-B/−4501MIK(a) oligonucleotide to abolish competition at the Ikaros (a) element is considered below.

Complexes D and E (hRFC-B/−4534; lane 12) were identified as involving binding of the Sp family of proteins to GC-box (b) by competition with a Sp1 consensus oligonucleotide (lane 14), perturbation of the DNA–protein complex by Sp1 antisera (lanes 16) and loss of competition after mutation of GC-box (b) [hRFC-B/−4534GC(b)] (lane 15). However, for the hRFC-B/−4501 oligonucleotide, including the previously characterized GC-box (a) [14], anomalous results were obtained since (i) the IBKS4 Ikaros consensus sequence completely abolished DNA binding for both complexes F and G (lane 20), yet (ii) Sp1 consensus sequence partially competed with complex F (lane 21) and (iii) Sp1 antisera supershifted most of complex F (lane 24). Furthermore, (iv) both Ikaros (a) and GC-box (a) mutants effectively competed for complexes F and G with Sp1 hRFC-B/−4501 probe but probably reflect the extensive overlap between the Ikaros (a) and GC-box (a) binding sites in this region (Figure 1B). For complex F, the abolition of DNA binding in the presence of IBKS4, and the lack of complete Sp1 competition, combined with the nearly complete supershift by Sp1 antisera, may partly reflect the presence of multiple DNA–protein complexes co-migrating as a single complex F and/or, possibly, protein–protein associations with Sp1. In spite of these ambiguities, the gel shift results with the hRFC-B/−4501 probe strongly suggest binding of Sp1 and Ikaros proteins to the overlapping Ikaros (a) and GC (a) elements in the −4501 to −4475 region.

Complex H (hRFC-B/−4434 oligonucleotide; lane 26) was identified as a USF protein bound to E-box (a) by competition with
Regulation of the human reduced folate carrier B promoter

Figure 2 In vitro binding of USF, Ikaros and Sp proteins to the hRFC-B promoter

Gel shift assays were performed with HT1080 nuclear extracts as described in the Materials and methods section. Double-stranded hRFC-B/-4573 (positions −4573 to −4554), hRFC-B/-4565 (−4565 to −4545), hRFC-B/-4561 (−4561 to −4475), hRFC-B/-4534 (−4534 to −4506) and hRFC-B/-4434 (−4434 to −4414) oligonucleotides were used (Figure 1B and Table 2). Results are shown for labelled probes without HT1080 nuclear extract (lanes 1, 6, 11, 17 and 25), and with 12 µg of extract (lanes 2, 7, 12, 18 and 26). Competitions (150-fold molar excess) included unlabelled hRFC-B/-4573 (lane 3), hRFC-B/-4565 (lane 8), hRFC-B/-4501 (lane 13), hRFC-B/-4501 (lane 19) and hRFC-B/-4434 (lane 27) oligonucleotides, an Ikaros consensus oligonucleotide (IKBS4; lanes 4, 9 and 20), an Sp consensus oligonucleotide (lanes 14 and 21), an E-box consensus oligonucleotide (lane 28), hRFC-B mutant oligonucleotides (lanes 5, 10, 15, 22, 23 and 29), Sp1 supershifts (lanes 16 and 24) and USF1 supershifts (lane 30). Complexes are indicated by upper-case letters and are described in the text. NS, non-specific complex; NE, nuclear extract; ab, antibody.

Figure 3 In vivo binding of Ikaros, Sp1, USF-1 and USF-2 to the hRFC-B promoter

HT1080 chromatin bound to transcription factor complexes were immunoprecipitated with Sp1, USF-1 and Ikaros antisera. Immunoprecipitations with IgG and no antibody were used as negative controls. PCR amplifications were performed as described in the Materials and methods section. Upper panel: the PCR products for promoter B (positions −4560 to −4353) are shown. Lower panel: a negative control for each immunoprecipitation amplified with coding exon-specific primers for the BST2 gene is shown. The amplicons were visualized on 2 % agarose gels. Ab, antibody; IK, Ikaros.

Mutagenesis of transcription factor binding elements in the hRFC-B promoter identified by gel shift assays

To assess further the functional importance of the major cis elements localized within the 158 bp hRFC-B core promoter (Figure 1B), full-length promoter constructs analogous to hRFC-B/-5422 were prepared containing mutations in the major cis elements identified on the gel shift assays (Table 2). With the exception of GC-box (a) and Ikaros (a), the cis element mutants (see above) significantly perturbed transcription factor binding in vitro. Promoter constructs including these same mutations in pGL3-basic were transfected into HT1080 cells, and luciferase activities were compared with that of the wt hRFC-B/-5422 promoter construct. As summarized in Figure 4, pronounced inhibitions of promoter activity were observed for mutations of both GC-boxes (a) and (b) (~80 % inhibition for both mutants) and E-box (a) (~90 % inhibition). Mutations of the Ikaros-binding sites resulted in variable and opposing effects on promoter activity, ranging from activation [Ikaros (c) and (a); ~15–30 % increase in luciferase activity] to inhibition [Ikaros (b); ~20 % decrease in luciferase activity]. Taken together, these results strongly support the notion that the cis elements depicted in Figure 1(B) and identified on gel shifts are functionally important in the transcriptional regulation of the hRFC-B core promoter.

Co-transfections of the hRFC-B promoter with Sp, USF and Ikaros expression constructs into Drosophila Mel-2 cells

The Sp and USF proteins are ubiquitously expressed transcription factors, yet are implicated in regulating tissue-specific or developmental expression of a number of genes [22–24]. Sp proteins...
transactivation by Sp1/USF2a, in contrast with the results of The effects of Ikaros 2 were partly reversed (was only slightly less (was only slightly less than 85%) repressive of other additions (IK2 and IK8 in Figure 5B). However, in the presence of the dominant-negative Ikaros 2 and 8 (each at 250 ng) were transcriptionally inert in the absence of high-affinity DNA binding, only Ikaros 1–3 significantly bind to DNA whereupon they can either activate or repress transcription by alternative splicing of exons 3–7 [19] as well as the related factors typified by Aiolos [26] and Helios [27,28]. Since at least three of the four internal zinc fingers (i.e. exons 3–5) are needed for high-affinity DNA binding, only Ikaros 1–3 significantly bind to DNA whereupon they can either activate or repress transcription [19,29]. Conversely, Ikaros 4–8 are dominant-negative isoforms that bind poorly to DNA and interfere with cellular response to the DNA-binding forms via formation of protein dimers [19,29].

To assess directly the role of USF proteins in hRFC-B transactivation and their relationships with the Sp response, Sp and USF cDNA constructs in pPac vector were co-transfected with the hRFC-B/−4590 construct [includes GC-boxes (a) and (b) and E-box (a); Figure 1B] into Drosophila Mel-2 cells that provide a null background for both these transcription factors [30]. Sp1, USF1 and USF2a, each individually, showed a dose-dependent stimulation of luciferase activity in Drosophila Mel-2 cells over the low level seen in empty pPacO (Figure 5A). Sp3 was significantly less stimulatory (1.5-fold over pPacO) of hRFC-B/−4590 promoter activity than Sp1 at an equivalent dose (results not shown). Of particular interest, the transactivating effects of Sp1 and USF1 or USF2a were highly synergistic (Figure 5B shows the combined effects of 25 ng of Sp1 with 500 ng of USF1 or USF2a).

Additional experiments were performed to assess the effects of Ikaros proteins on hRFC-B/−4590 promoter activity. Both Ikaros 2 and 8 (each at 250 ng) were transcriptionally inert in the absence of other additions (IK2 and IK8 in Figure 5B). However, in combination with Sp1 and USF1, Ikaros 2 was notably repressive, resulting in an approximate 90% inhibition of promoter activity. The effects of Ikaros 2 were partly reversed (~3-fold) in the presence of the dominant-negative Ikaros 8. Whereas Ikaros 2 was only slightly less (~85%) repressive of hRFC-B/−4590 transactivation by Sp1/USF2a, in contrast with the results of Sp1/USF1, the effects of Ikaros 2 were nearly completely reversed by Ikaros 8.

These results clearly demonstrate that Sp and USF proteins, individually, are capable of stimulating the 158 bp hRFC-B core promoter. However, both Sp1 and USF proteins are required for high levels of promoter transactivation. In addition, our results strongly imply that hRFC-B transactivation can be significantly modulated by the presence of the different Ikaros family members, including both DNA-binding and transdominant-negative forms, and that the magnitude of these effects differs slightly for Sp1, USF1, USF2a, Ikaros 2 (IK2) and Ikaros 8 (IK8). All transfections included a total of 500 ng of total vector including expression construct and empty pPacO. Cells were harvested after 24 h for luciferase assays. Luciferase activities were normalized to cellular protein contents. Results are expressed as means ± S.E.M. for three separate experiments.

Effects of Sp1, USF1 and Ikaros 2 transfections on hRFC-B transcript levels and promoter B activity in HT1080 cells

The results in Figure 5 with Drosophila Mel-2 cells co-transfected with the hRFC-B promoter and transcription factor expression constructs provide compelling evidence that both Sp1 and USF1
are each capable of stimulating the core promoter and, in combination, could elicit a highly synergistic transactivation response. Moreover, Ikaros 2 potently repressed the high-level promoter activity achieved with USF1 and Sp1, although the effects of Ikaros 2 were partially reversed by the dominant-negative Ikaros 8.

An important extension of these experiments involves the effects of these transcription factors on levels of endogenous hRFC transcripts transcribed from non-coding exon B in HT1080 cells, as a direct measure of endogenous hRFC-B promoter activity. To address this question, HT1080 cells were transiently transfected with expression constructs (in pCDNA3) for Sp1, USF1 or Ikaros 2, and after 48 h, hRFC transcripts with B 5′-UTR sequence were assayed by real-time RT–PCR with exon B-specific primers. In three experiments, both USF1 and Sp1 treatments resulted in significant and reproducible (2.2- and 1.9-fold respectively) increases in the levels of endogenous hRFC-B transcripts, whereas Ikaros 2 was repressive (∼20%) (Figure 6A). Thus the effects of these exogenous transcription factors on the endogenous hRFC-B promoter in HT1080 cells parallel the changes in hRFC-B core promoter activity in Drosophila cells transfected with the hRFC-B/−4590 reporter gene construct. Similarly, cotransfection of HT1080 cells with USF1 and Sp1 stimulated (not shown) and Ikaros 2 suppressed (Figure 6B) hRFC-B/−4590 promoter activity (Figure 6B). The repressive effects of Ikaros 2 were completely reversed in co-transfections with Ikaros 8.

DNA-binding Ikaros proteins have been reported to achieve at least part of their transcriptional effects via interactions with chromatin remodelling complexes, by recruiting the co-repressors, Mi-2β, Sin3A and Sin3B and the class I HDACs (histone deacetylases) 1 and 2 [31,32]. If such a mechanism were operative for hRFC, loss of hRFC-B transcripts and promoter activity would be accompanied by decreased levels of acetylated chromatin associated with the B promoter. To assess the changes in acetyl histone H3 in HT1080 cells treated with Ikaros 2, we used ChIP with anti-acetyl histone H3 antibody (Lys-14). Chromatin was immunoprecipitated with IgG or acetyl H3 antibody and PCR-amplified in the linear range with primer spanning the hRFC-B core promoter region (positions −4560 to −3353). Although input signals for hRFC-B were comparable between the mock and Ikaros 2 transfectants, the level of acetyl histone H3 associated with the hRFC-B promoter decreased after treatment with Ikaros 2 (Figure 6C). As shown in the bottom panel, there was a 1.8-fold decrease in the level of acetylated histone H3, when immunoprecipitates were analysed over a 6-fold range of concentrations. No signals were detected in the absence of acetyl H3 antibody or after immunoprecipitation with IgG. For BST2 gene control amplifications, other than the input, no signals were detected. These results strongly suggest that transfections of HT1080 cells with Ikaros 2 result in significantly decreased acetyl histone H3 associated with the endogenous hRFC-B promoter accompanying decreased hRFC-B transcripts.

**DISCUSSION**

We previously localized the minimal promoter for hRFC-B to within 46 bp (positions −4493 to −4448) and demonstrated an important regulatory role for the Sp family of transcription factors in binding to a critical GC-box element within this region [14]. In the present study, we characterize the probable contributions of other cis elements and transcription factors, and document a potential role for chromatin remodelling in regulating promoter B activity.
Important regions involved in regulating promoter B were established by 5'- and 3'-deletions and transient transfections of HT1080 cells. From these results, oligonucleotide probes spanning a 158 bp core promoter region were designed and used in gel shift assays with HT1080 nuclear extracts to identify binding sites for key transcription factors. In vivo binding of transcription factors identified on gel shifts was confirmed by ChIP.

The important regulatory role previously described for the GC-box (a) element [14] was confirmed on gel shifts and by reporter gene assays of a GC-box (a) mutant construct. Furthermore, additional bound Sp proteins were localized to another GC-box, designated GC-box (b), and USF proteins were bound to a non-canonical E-box (a), flanking GC-box (a). When GC-box (b) and E-box (a) were individually mutated so as to impede their in vitro protein binding, promoter activity was inhibited.

Given the close proximity of the hRFC-B transcription start sites to these GC- and E-box elements, the notion of an essential protein binding, promoter activity was inhibited. 

A regulatory role for the family of Ikaros proteins with hRFC-B was also suggested by our studies, via binding to three Ikaros binding sites [Ikaros (a)–(c)] upstream of GC-box (a). The functional importance of these elements was further implied by transient transfections of the Ikaros (a), (b) and (c) mutant constructs into HT1080 cells. In Drosophila Mel-2 cells, either Sp1 or USF transcription factors could transactivate the hRFC-B−/− 4590 reporter construct, although maximal transactivation required both Sp1 and USF1. USF2a was also capable of transactivating the hRFC-B promoter; however, Sp3 was largely inert. Both USF1 and Sp1 stimulated hRFC-B+/+ 4590 reporter activity in HT1080 cells (not shown) and increased the levels of endogenous hRFC-B transcripts in transiently transfected HT1080 cells. Co-operative interactions between Sp1 and USF proteins were previously described for the human transcobalamin II [33], cystathionine-β-synthase 1b [34] and, most recently, deoxycytidine kinase [20] promoters. With deoxycytidine kinase, Sp1 and USF1 were co-immunoprecipitated [20], thus providing direct evidence for a physical interaction between these proteins. The synergistic stimulation of the hRFC-B promoter in Drosophila Mel-2 cells with Sp1 and USF1/2a suggests that both Sp and USF proteins are essential for high-level transactivation. Accordingly, the net level of hRFC-B promoter activity achieved would probably reflect relative levels of active Sp1, Sp3, USF1 and USF2a transcription factors within cells.

A regulatory role for the family of Ikaros proteins with hRFC-B was also suggested by our studies, via binding to three Ikaros binding sites [Ikaros (a)–(c)] upstream of GC-box (a). The functional importance of these elements was further implied by transient transfections of the Ikaros (a), (b) and (c) mutant constructs into HT1080 cells. In Drosophila Mel-2 cells, co-transfection with Ikaros (a), (b) and (c) mutant constructs into HT1080 cells. In Drosophila Mel-2 cells, co-transfection with Ikaros 2 repressed hRFC-B transactivation by Sp1 and USF1. Whereas the dominant-negative Ikaros 8 was itself transcriptionally inert, this form reversed the repressive effects of Ikaros 2. The extent of this reversal was different in the presence of USF1 versus USF2a. Ikaros 2 also decreased the endogenous level of hRFC-B transcripts in HT1080 cells, consistent with its repressive effects on hRFC-B reporter gene activity.

T-cell malignancies in mice result from a loss of Ikaros expression or overexpression of dominant-negative Ikaros isoforms [35]. In childhood ALL (both B-precursor and T-ALL), expression of the dominant-negative Ikaros isoforms (Ikaros 4, 6–8) was also detected at high frequencies [36,37]. This provides at least a partial explanation for the extraordinarily wide range of hRFC expression previously reported in ALL lymphoblasts [11]. However, any effect of Ikaros proteins on hRFC-B promoter activity in ALL would probably be compounded by those of other transcription factors, such as the USF and the Sp families of proteins.

DNA binding by the family of Ikaros proteins has been associated both with promoter transactivation and repression [19,29].

Ikaros has been reported to associate with the SWI/SNF chromatin remodelling complex [31] and localize with inactive genes in centromeric heterochromatin [27]. Repression by DNA-binding Ikaros isoforms has been shown to occur via association with the CtBP co-repressor [38] or by association with chromatin remodelling activities such as Mi-2β and the Sin3A and Sin3B co-repressor complexes including HDACs [31,32]. It is probable that the levels and activities of the various binding partners for Ikaros determine the nature of the complexes that form and whether they repress or activate transcription. When chromatin was prepared from HT1080 cells treated with Ikaros 2 under conditions that repress hRFC-B reporter gene activity and endogenous hRFC-B transcripts, and analysed by ChIP with anti-acetyl histone H3, levels of acetylated histone H3 were decreased compared with mock-transfected cells. Thus the repressive effects of Ikaros 2 on hRFC-B transcription would seem at least in part to be reflected in decreased acetylation of the histones associated with the −4560 to −4353 core promoter region.

Interestingly, treatment of the Mtx-resistant MDA-MB-231 breast cancer subline with the HDAC inhibitor, trichostatin A, was previously reported to have no net effect on total levels of hRFC transcripts, even though loss of hRFC expression was associated with decreased acetylation of histones H3 and H4 [39]. However, these results must now be interpreted in terms of a multiplicity of hRFC promoters and 5′-UTRs [15] whereby the effects of trichostatin A may be restricted only to a limited number of 5′-UTRs/promoters with no significant effect on other hRFC 5′-UTRs/promoters. Thus the recruitment of chromatin remodelling machinery by specific transcription factors or drug treatments could represent a potentially selective mechanism for modulating hRFC 5′-UTR/promoter usage. This will be tested in future studies.

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