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Runx1 binds as a dimeric complex to overlapping Runx1 sites within a palindromic element in the human GM-CSF enhancer

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ABSTRACT

Runx1 is a developmentally regulated transcription factor that is essential for haemopoiesis. Runx1 can bind as a monomer to the core consensus sequence TGTGG, but binds more efficiently as a hetero-dimer together with the non-DNA binding protein CBFβ as a complex termed core binding factor (CBF). Here, we demonstrated that CBF can also assemble as a dimeric complex on two overlapping Runx1 sites within the palindromic sequence TGTGGCTGCCCA in the human granulocyte macrophage colony-stimulating factor enhancer. Furthermore, we demonstrated that binding of Runx1 to the enhancer is rigidly controlled at the level of chromatin accessibility, and is dependent upon prior induction of NFAT and AP-1, which disrupt a positioned nucleosome in this region. We employed in vivo footprinting to demonstrate that, upon activation of the enhancer, both sites are efficiently occupied. In vitro binding assays confirmed that two CBF complexes can bind this site simultaneously, and transfection assays demonstrated that both sites contribute significantly to enhancer function. Computer modelling based on the Runx1/CBFβ/DNA crystal structure further revealed that two molecules of CBF could potentially bind to this class of palindromic sequence as a dimeric complex in a conformation whereby both Runx1 and CBFβ within the two CBF complexes are closely aligned.

INTRODUCTION

Runx1 (also known as AML1, CBFα2 or PEBP2αB) is a Runt-domain transcription factor (1,2) that is essential for haemopoiesis (3,4). Runx1 is a member of a conserved family of closely related proteins that include mammalian Runx1, Runx2 and Runx3, and the Drosophila protein Runt (1,2). Runx1 binds to TGTGGNNN core sequences, typically TGTGGTTT or TGTGGTCA, as a heterodimer of Runx1 and CBFβ, termed core binding factor (CBF) (2,5,6). Although CBFβ does not directly contact DNA, it helps to stabilize Runx1 binding (2). While TGTGGTT is the most commonly observed natural CBF-binding sequence, in vitro studies reveal that TGCGGT is also a high affinity, but much less often encountered, CBF-binding sequence (2,5,7).

Runx1 binds to DNA via the Runt domain, which shares some similarity with the Rel domain of NF-κB/Rel family proteins that also recognize GG core sequences (8,9). However, NF-κB binds as an obligate dimer that employs two Rel family proteins to bind to palindromic sequences such as GGGAAATTCCC (10). This is in contrast to Runx1 which interacts efficiently with single TGTGGNNN consensus motifs (6). The crystal structure of a Runx1-Runt domain/CBFβ complex bound to the DNA sequence TGCGGT TG has been determined, revealing contacts with both the bases and the phosphate...
backbone throughout this 8-bp sequence (11). Similar results were obtained by a parallel study of the Runx1 binding site TGTGGTTG (12). Runx1 clamps the phosphate backbone between the major and minor groove, while forming Rel-like base-specific contacts with the GG sequence in the major groove (11). In this way, Runx1 utilizes multiple protein:DNA contacts to enable efficient binding to single sites (11).

Runx1 synerizes with a variety of other factors to regulate composite elements. For example, Runx1 and Ets-1 interact directly and bind cooperatively to enhancers within the TCRα and TCRβ loci (13,14). Runx1 and C/EBP family proteins synerize in the activation of adjacent binding sites within the M-CSF receptor gene promoter (15). Runx1 and c-Myb synerize in the activation of adjacent sites in the TCR-δ enhancer, although in this instance synergy does not depend on cooperative binding (16). Runx1 also possesses an interaction domain within the C terminal region that mediates homodimerization, and this may promote binding to regulatory elements containing multiple Runx1 binding sites (17). The homodimerization domain appears to be distinct from the region of Runx1 required for interaction with other factors, such as Ets-1 (14). This domain is less structured than the Runt domain, and no detailed determination of its structure is available.

Runx1 shares some similarities with another class of Rel domain transcription factor, the NFAT family of proteins, which bind to GGAAANN consensus sequences. Like Runx1, NFAT uses multiple interactions to bind to DNA efficiently as a monomer. NFAT uses a single Rel domain to bind in the major groove to an essential GGA core sequence, and also has contacts in the minor groove along the next 4-bp downstream of the GGA core (18). NFAT typically functions and binds cooperatively together with other transcription factors such as AP-1 (18,19). However, NFAT family proteins can also bind as homodimers to NF-kB-like sequences conforming to the consensus sequence GGAAATTC (18,20,21). This raises the possibility that Runx1, like NFAT, might also be able to bind to palindromic sequences containing two overlapping Runx1 binding sites. However, to date we are not aware of any reports of such examples.

The human granulocyte macrophage colony-stimulating factor (GM-CSF or CSF2) gene is a key target of regulation by Runx1 (22–24). GM-CSF is a pro-inflammatory cytokine produced by activated T cells and mast cells that is induced by stimuli that activate the immune system. The expression of the human GM-CSF gene is under the control of a promoter region located just upstream of the transcription start site and an inducible enhancer located 3-kb upstream, which are both activated by kinase and calcium signalling pathays that in T cells are activated via the T-cell receptor (TCR) (19,25–27). We and others have previously demonstrated the presence of functional Runx1 sites in both the GM-CSF promoter (22–24) and the inducible enhancer (22). GM-CSF gene expression can also be inhibited by the repressive AML1/ETO fusion protein found in t(8;21) chromosomal translocations in acute myeloid leukaemia (23,24). These translocations delete the homo-dimerization domain, and the addition of the ETO domain converts Runx1 from an activator to a repressor.

The GM-CSF enhancer is defined as a 717-bp BglIII fragment of DNA that requires interactions with multiple transcription factor sites for function (25,28). Within the GM-CSF enhancer, Runx1 binds to the GM450 element within a region that is normally occluded by a positioned nucleosome (Figure 1A; 28,29). Activation of the enhancer by TCR signalling pathways results in recruitment of NFAT and AP-1 to sites located within this same nucleosome (N2) and to additional sites in an adjacent nucleosome (N1), leading to the rapid eviction of both nucleosomes (summarized in Figure 1A). This process results in the rapid creation of a DNase I hypersensitive site (DHS), a substantial increase in chromatin accessibility, and the recruitment of Runx1 to a site that was previously unoccupied. This mechanism thereby allows constitutively expressed factors such as Runx1 and Sp1 to bind in a highly inducible fashion.

In this study, we demonstrated that the GM-CSF enhancer GM450 Runx1 binding site TGTGGGTTG ACA actually has two overlapping TGTGGNNN Runx1-like elements where the TGTGG core sequences (underlined) exist 4 bp apart within this palindromic sequence. Furthermore, we obtained evidence that both sites become occupied in activated T cells and mast cells, and demonstrated that both sites contribute to enhancer function. Computer modelling predicted that the palindromic element is able to accommodate binding of two Runx1/CBFβ complexes closely aligned on the same face of the DNA helix to form multiple close associations along an axis perpendicular to the DNA.

MATERIALS AND METHODS

Cell culture

The CEM and Jurkat human T-cell lines were cultured in RPMI (Invitrogen) supplemented with 10% FCS, 4 g/l D-glucose, 1 mM sodium pyruvate, 1 mM MEM essential amino acids (Invitrogen), 1 × MEM non-essential amino acids (Invitrogen), 100 U/ml penicillin and 100 μg/ml streptomycin. C42 human IL-3/GM-CSF transgenic mouse T lymphoblasts and mast cells were prepared and cultured as described previously (29,30).

In vivo footprinting analyses

Footprinting analyses were performed essentially as described (31,32). C42 transgenic mouse T lymphoblasts were either stimulated for 4h with 20 ng/μl phorbol 12-myristate 13-acetate (PMA) and 1 μM calcium ionophore A23187, or left unstimulated, before treatment with 0.2% dimethyl sulphate (DMS) in phosphate buffered saline (PBS) for 5 min at room temperature. Specific DNA cleavage sites were detected using ligation-mediated (LM)-PCR as described (31,32). These analyses were performed with two sets of 3 nested primers designed to detect DNA cleavage sites on either the upper or the lower strand of the GM420/GM450 region of the human GM-CSF enhancer, plus the linker primer.
The three primers within each set are designated EB for the biotinylated primer used at the first stage, EP for the primer used at the second PCR stage and EL for those used in the final labelling stage. The 50 primers used to detect cleavage on the lower strand (plus their annealing temperatures shown in brackets) were EB3F2, CACAGC CCATCGGAGC (52°C/C14°C); EP3F20, CTGAGTCAGCAT GGCTGGC (62°C/C14°C); EL3F, GCATGGCTGGCTATCGGT TGACACTG (68°C/C14°C). The 30 primers used to detect cleavage on the upper strand (plus their annealing temperatures) were EB2R2, GCCCAAGTCAGCACAAAC (56°C/C14°C); EP2R, GTCAGCACAAACAGGACAGAAATC (64°C/C14°C); EL2R20, AGGACAGAAATCCATGGGT TGACACTG (64°C/C14°C).

Electrophoretic mobility shift assays

To study binding of purified CBF, we obtained a sample of the purified recombinant Runx1/CBFβ complexes that had been used previously by Bravo et al. (11) to prepare crystals of CBF bound to DNA. These complexes contained just the His-tagged Runt domain region of Runx1 (residues 50–183), and residues 2–135 of CBFβ. To study binding of native CBF complexes we prepared nuclear extracts from unstimulated Jurkat T cells, essentially as described previously (25).

Double stranded oligonucleotides were labelled by end-filling with 32P-dCTP plus unlabelled dATP, dGTP and dTTP, and the labelled probes were purified by polyacrylamide gel electrophoresis of LM-PCR products. Open circles indicate protected bases, and filled circles indicate hyper-reactive bases. Bases are numbered relative to the upstream BglII site. (C) Quantification of the in vivo protection of the CBF site on the lower DNA strand. These densitometric traces of lanes 9 (nil) and 10 (PMA/I) in Figure 1B reveal DMS reactivity in mast cells before and after stimulation.

Figure 1. Transcription factors and nucleosomes associated with the human GM-CSF enhancer. (A) Map of the 717 bp BglII fragment of the enhancer showing the previously determined locations of positioned nucleosomes (N0–N3) and transcription factors bound to the GM-CSF enhancer in T cells before and after stimulation with PMA and calcium ionophore. The composite NFAT/AP-1 elements are termed GM330 and GM420, and the Runx1 element is termed GM450. The dotted ovals signify nucleosomes that become mobilized after stimulation which do not reside in fixed positions. (B) In vivo DMS footprints within the GM-CSF enhancer. Cultured C42 transgenic mouse T cells and mast cells were assayed before (−) and after (+) stimulation for 4 h with PMA and calcium ionophore. Either genomic DNA (G) or intact cells were treated with DMS, and the modified bases were cleaved with piperidine and identified by polyacrylamide gel electrophoresis of LM-PCR products. Open circles indicate protected bases, and filled circles indicate hyper-reactive bases. Bases are numbered relative to the upstream BglII site. (C) Quantification of the in vivo protection of the CBF site on the lower DNA strand. These densitometric traces of lanes 9 (nil) and 10 (PMA/I) in Figure 1B reveal DMS reactivity in mast cells before and after stimulation.
0.25–2.0 pmol of CBF with 0.2 ng of labelled probe and 100 ng of poly(dI·dC) in a 16 μl reaction containing 10% glycerol, 20 mM HEPES pH 7.9, 50 mM NaCl, 80 mM KCl, 3 mM MgCl₂, 1% DTT, 0.1 mM PMSF, 5 μg/ml aprotinin, 5 μg/ml leupeptin, 0.1 mg/ml BSA at room temperature for 10 min and 10 min on ice before analysis on 4% polyacrylamide gels run at 4°C.

The oligonucleotides used to prepare DNA probes had the following sequences, which include a 2 base 5' overhang at each end:

- TCR-δ Runx1, AGGCATGTGGTTTCCAACCGTT and TGAACGGTTGGAACACATGC;
- Ideal, AGATGTGTGGTTAACCACAAAC and TGAACGGTTGGAAACCACATGC;
- GM450 and AGGTTTGTGGGCAGCCACACAT;
- WT GM450, AGATGTGTGGCTGCCCACAAAC and AGGTTTGCGGTCGACCGCACAT;
- Crystal, AGATGTGCGGTCGACCGCAAAC and AGGTTTGTCCTTAACCACACAT.

 Autoradiography and densitometry

Images were collected on a BioRad Phoros FX Plus phosphorimager. Densitometry and band quantitation was performed using BioRad Quantity One software.

**Plasmid construction**

The previously described constructs pGM and pGM-GME contain the −627 to +28 region of the human GM-CSF promoter (GM627) in the absence or presence of a 717 bp Bgl II fragment of the human GM-CSF enhancer, respectively (33), inserted into the firefly luciferase reporter gene plasmid pXPG which includes highly efficient transcription termination and polyadenylation elements upstream of the enhancer (34).

Site-directed mutagenesis was performed as previously described (28) to change the TGTGG core sequences located at positions 443 and 452 in the GM-CSF promoter (GM627) in the absence or presence of a HindIII–Xba I of the Renilla luciferase gene excised from the control plasmid pRL-TK (Promega). This vector retains the same configuration of vector backbone, upstream SV40 polyA/termination sites, and GM-CSF enhancer and promoter as in pGM-GME, and is designed to respond in the same way upon transfection and stimulation.

**Transient transfection and luciferase assays**

Aliquots of 4.5 × 10⁶ Jurkat T cells were transfected by electroporation with plasmid DNA purified by two rounds of CsCl gradient centrifugation. Cells were transfected with 5 μg of Firefly luciferase plasmid plus 1 μg of the control pXRL-GME Renilla luciferase plasmid. Cells were cultured for ~21 h post-transfection, before stimulation with 20 ng/ml PMA and 2 μM calcium ionophore A23187 for 8 h. Cells were then washed in PBS and assayed for luciferase reporter gene activities using the Promega dual luciferase assay kit and a Berthold LB-940 microplate luminometer. Data were collected from at least two independently prepared clones of each construct, which were in each case found to behave in like fashion. A minimum of 6, and up to 21 independent transfections were performed for each construct.

**Computational modelling of the dimeric form of Runx1-CBFβ-DNA ternary complex**

An extended and detailed explanation of the computational modelling and refinement of complex can be found in the Supplementary Methods. Briefly, the Run1-CBFb-DNA coordinates defined in Bravo et al. (11, 35) were superimposed over a model of guide B-DNA containing two Runx1 DNA binding sites generated using the nucleic acid builder tool (36) in the AmberTool package and the 21 bp GM-CSF enhancer sequence ATGTTGGGCCGCATGGAGATT. The crystal structure of the human Runx1-CBFβ-DNA ternary complex (11) was used as template to model two Runx1–CBFβ complexes bound to the guide B-DNA using MODELLER (37). Sequences and templates were aligned so that DNA molecules of the template were aligned to the corresponding DNA binding sites in the guide B-DNA preserving the native Run1–CBFβ–DNA interactions described in the crystal structure (11). The refinement of the model consisted of two rounds of energy minimization and a short molecular dynamic simulation of the solvated complex in order to ensure a reasonable stereochemical geometry, resolve steric hindrances, and assess the stability of the complex. The energy minimization and molecular dynamic simulation was performed in AMBER 10 (38) using the ff99SB force field (39), as follows: On the first round of minimization, protein and DNA atoms were kept restrained and solvent was relaxed. On the second round, restraints on DNA and protein atoms, with the exception of those mediating the interactions between protein and DNA, were lifted and system was further minimized. The temperature of the system was then raised gradually to reach 300K, while remaining restraints were also gradually removed during the heating process. Finally, the system was equilibrated for 0.5 ns and simulated for 4 ns. The final model was obtained after clustering of the entire molecular dynamic simulation, selecting a representative on each cluster, and assessing the quality of each representative using PROCHECK (40). The convergence of the simulation and stability of the complex was ensured by plotting the root mean square deviation and total energy as a function of the simulation time.
RESULTS
Runx1 binds to 2 overlapping sites within the GM-CSF enhancer in vivo upon stimulation

We previously defined the GM450 Runx1 binding site TG CCCACA that has a TGTGG core sequence located at position 452 within the human GM-CSF enhancer (22). This site forms specific Runx1/CBFβ complexes as previously defined with the aid of specific antibodies and DNA competitors, and it can be used to functionally replace the Runx1 site located within the GM–CSF promoter (22). To study the regulation of the human GM-CSF locus in defined populations of normal cells, we employed the transgenic mouse line C42 containing the intact human GM-CSF locus on a 130 kb BAC segment of DNA (30).

To study Runx1 interactions with the GM-CSF enhancer inside live cells we employed DMS in vivo footprinting of T cells and mast cells prepared from C42 mice (29) (Figure 1B). With this assay we can identify sites of in vivo protein:DNA interactions by treating cells with dimethyl sulphate (DMS), which modifies any G bases that are not occupied by interacting factors. To this end, cells were treated with DMS before (−) or after (+) stimulation for 4 hours with 20 ng/ml PMA and 2 μM calcium ionophore A23187. As a control for protein-free DNA, we also treated purified genomic DNA with DMS (G).

Modified bases were cleaved by incubation with piperidine, and the specific cleavage sites were detected by ligation-mediated PCR. Figure 1B displays the results obtained after analysis of sites on both the upper and the lower strands of DNA. The results are summarized below with protected bases marked by open circles, and hypersensitive bases marked with filled circles. Before stimulation, there was no evidence for any interactions between any factors and the enhancer, as the patterns were essentially the same as those obtained by treating genomic DNA with DMS. After stimulation, there was clear evidence of efficient binding of factors to the Runx1, NFAT and AP-1 sites within this region of the enhancer. Furthermore, the Runx1 site supported not just one, but two distinct sets of footprints. In addition to binding at the previously defined TGTGG motif at position 452 (lower strand, lanes 8 and 10), there was strong protection of the Runx1-like element TGTGGCT G located at position 443 (upper strand, lanes 3 and 5). There was also a hypersensitive reaction with a G at position 451 on the lower strand, which is symptomatic of an altered DNA conformation induced by factor binding.

Particularly in the mast cells, it was evident that there was very strong protection of both Runx1 sites (lanes 5 and 10), indicating that both sites are likely to be stably occupied simultaneously in a substantial proportion of cells. Quantitation of a densitometric analysis of the data in lane 10 (for the lower strand) in stimulated mast cells (Figure 1C) indicated that only about 16% of all G bases at position 455 within the motif at position 452 remained accessible. A similar densitometric analysis of lane 5 (the upper strand) indicated that only 34% of G bases at positions 444 and 446 within the motif at position 443 remained accessible after stimulation (data not shown). Based on this, we can calculate that 66% of all motifs at position 443 on the upper strand become occupied. If only 16% of motifs at position 452 on the lower strand remain free, then this implies that a minimum of 50% (and maximum of 66%) of all GM450 palindromic Runx1 sites must be stably occupied at both sites in stimulated mast cells.

The GM-CSF enhancer Runx1 element can recruit two CBF complexes at the same time

We used EMSAs to study the in vitro binding of recombinant CBF complexes to the Runx1 sites (Figure 2A and B). For this purpose, we obtained a sample of the original purified recombinant CBF complex that had been used previously by Bravo et al to determine the crystal structure of the CBF/DNA complex (11). This recombinant CBF contained the His-tagged Runt domain region of Runx1 (residues 50–183), and the region of CBFβ (residues 2–135) that mediates interactions with Runx1. With 0.25 pmol of CBF (lane 1), two different mobility complexes were observed with the wild type probe (WT GM450). Although we have been unable to determine why two different mobility complexes are detected, it is likely that they both contain a single molecule of Runx1, because both complexes were still detected when either the position 443 or 452 Runx1 sites were mutated individually (lanes 5 and 8). These complexes were both regarded as being specific for the TGTGG motifs, because no binding was detected when both Runx1 sites were mutated simultaneously (lanes 11–13). It is possible that the lower band represents a Runx1–DNA complex, from which CBFβ has dissociated, but we have been unable to confirm this.

When amounts of up to 2 pmol of recombinant CBF were added to the intact GM450 Runx1 probe, increasing amounts of a third lower mobility complex were observed (lanes 2–4). This upper complex was assumed to be a dimeric CBF complex containing two Runx1/CBFβ heterodimers, because it was not detected when either individual Runx1 site was mutated (lanes 5–10, Figure 2B).

In one published study, the structure of the CBF/DNA complex was determined using the DNA sequence TGG GTTG. To determine whether we could use this structure as a basis for constructing a computer model of a dimeric complex, we designed the palindrome TGCGGTCGACC ACA that matches the ideal consensus TGTGGTCA or TGTGGTTA ACC ACA that matches the ideal consensus TGTGGTCA on
Figure 2. EMSAs of the GM450 Runx1 binding site. (A) Sequences of the wild-type and mutated GM450 Runx1-binding probes, plus the Ideal, the IdealΔ1 and the crystal structure-based palindrome sequence used in (B–G), and the TCRδ single Runx1 site. The Crystal sequence is a palindrome derived from the DNA sequence used to prepare crystals by Bravo et al. (11). The Ideal sequence is a palindrome derived from the ideal Runx1 consensus binding site, and the IdealΔ1 sequence has the second Runx1 site in this element mutated. The 8 bp regions spanned by the two Runx1 binding sites are indicated by bars above the sequence, and the mutated GG core elements are shown as underlined in bold. Bases in lower case represent 2 base 5’ overhangs at each end that become double stranded when probes are labelled. (B) EMSAs with increasing concentrations recombinant Runx1/CBFβ complexes binding to the wild-type and mutated GM450 sequences and the Crystal sequence. Bands corresponding to a monomer and a dimer of Runx1 are marked. (C and D) EMSAs with 6 µg Jurkat cell nuclear extract with (C) the wild-type GM450 (lane 1), the mutated GM450Δ443 (lane 2) and the crystal (lane 3) sequences and (D) the ideal sequence (lane 1) in the presence or absence of an antibody against either the N-terminal (Runx-N) (lane 2) or C-terminal (Runx-C) (lane 3) of Runx1, or control IgG (lane 4). (E) Competition assay with 6 µg of Jurkat nuclear extract bound to labelled Ideal probe (lane 1) and competed with 25 ng of either Ideal (lane 2), TCRδ (lane 3) or GM450Δ443Δ452 (lane 4) unlabelled oligonucleotides. (F) EMSAs with 6 µg Jurkat cell nuclear extract bound to the Ideal probe (lanes 1 and 7) and competed with increasing amounts of either unlabelled ideal (lanes 2–6) or IdealΔ1 (lanes 8–12) oligonucleotides. (G) Densitometric quantification of the EMSAs in (F) showing the proportion of the monomeric and dimeric complexes remaining in the presence of competing Ideal or mutated IdealΔ1 oligonucleotides.
both strands (designated as Ideal in Figures 2A). Both the
GM450 and Crystal probes supported the binding of what
appeared to be monomeric and dimeric CBF complexes
(Figure 2C). This conclusion was supported by the fact
that a mutation of the 443 site led to disappearance of
the lower mobility complex.

The Ideal palindromic Runx1 consensus probe also
formed two distinct complexes that were assumed to be
CBF monomer and dimer complexes (lane 1 in Figure 2D
and E). The designated CBF monomer and dimer
complexes were both confirmed as specific Runx1
complexes because they (i) were both super-shifted by
two different Runx1 antibodies (lanes 2 and 3, Figure 2D),
but not by control IgG (lane 4, Figure 2D), and (ii) were both inhibited by oligonucleotides containing
the position 443 and 452 TGTGG motifs, plus the plasmid
pGM containing just the GM-CSF promoter (26). To
control for both transfection and stimulation efficiency
at the same time, we co-transfected cells with the
plasmid pXRL-GME. This plasmid was created from
pGM-GME by excising the firefly luciferase gene and
replacing it with the Renilla luciferase gene.

Firefly and Renilla luciferase plasmids were
co-transfected into cells that were then cultured for
~21 h, before stimulating the cells for 8 hours with
20 ng/ml PMA and 2 μM calcium ionophore A23187,
and harvesting of extracts for dual luciferase assays. We
first confirmed that there was no unwanted cross-
contamination of activities between firefly and Renilla
luciferase assays by transfecting Jurkat cells with each
plasmid alone and performing dual luciferase assays
(data not shown). We next demonstrated that both
the pGM-GME and pXRL-GME plasmids were induced by
a factor of ~100-fold upon stimulation (data not shown)
and showed that inclusion of the enhancer increased pGM
promoter activity by about 5 to 10 fold in CEM and
Jurkat cells (Figure 3). The introduction of single GG to
CC mutations within the position 443 or 452 motifs led to
a ~3-fold reduction of activity in CEM cells and to a
~2-fold reduction of activity in Jurkat cells (Figure 3).
The double mutation of both the position 443 and 452
TGTGG motifs essentially abolished enhancer activity in
CEM cells, and reduced activity of the enhancer in Jurkat
cells to ~2-fold the activity of the promoter alone (Figure 3). Hence, we have confirmed that both
Runx1 sites are required for efficient inducible enhancer function.

Computational modelling of a dimeric CBF complex
on a palindromic sequence

To determine (i) how overlapping Runx1 motifs might
accommodate two CBF complexes, and (ii) whether
dimer formation is potentially supported by interactions
between the two CBF complexes, we generated a
computer model of the dimer incorporating the Crystal
palindromic defined in Figure 2. For this purpose, we
mapped the Runx1-CBFβ coordinates determined by
Bravo et al for the single Runx1-CBFβ-DNA complex
(11) onto the two Runx1 motifs within GM-CSF
enhancer sequence ATGTGTTGGCTGCCCACAAAAC.
The core of the DNA binding regions, and in particular
the GG core elements, were aligned to those in the palin-
dromic sequence in order to preserve the geometry of the
interaction between Runx1 and DNA. Two major assump-
tions were made when modelling the structure of the
dimeric CBF complex: first, that each individual
CBF complex will recognize the DNA binding region
comparably to the monomeric form as reported by
Bravo et al., and second, that each individual CBF
complex would not undergo major conformation
changes when forming the dimeric CBF complex.

The raw structural model of the dimeric CBF complex
presented some minor steric clashes between the two CBF
complexes. In particular, some atoms of residues located
in the interface between the CBFβ subunits and the βE–F
loop [as reported in Bravo et al. paper (11)] were at an atomic distance inferior to the sum of van der Waals radii (data not shown). The reasons for this are because (i) during the modelling of the complex, the DNA molecule containing the palindromic sequences was modelled as straight B-DNA configuration, and was kept rigid, and (ii) the interactions between the Runx1 domains and DNA was maintained in the same conformation as in the crystal structure (11). However, when the complex was subjected to an energy minimization step and a short dynamic simulation, i.e. protein and DNA atoms were allowed to relax and change conformation, these steric clashes were largely resolved. PROCHECK did not report any major stereochemical problems in the minimized complex. Furthermore, the model of the complex remained stable throughout the simulation and the total energy of the complex converged to a minimum. From the simulation, we were able to derive several clusters of structures, and select representative examples of each. From these, we selected the structure of the complex that provided the best G-factor in PROCHECK, and this is presented in Supplementary Data File 1.

As represented in the model of this structure in Figure 4A, two CBF complexes can comfortably bind simultaneously to the DNA binding sites without any major steric impediment. The two GG core sequences of the DNA motifs are separated by one half turn of helix and exist in opposite orientations. This means that each Runx1 domain is rotated 180 degrees relative to the other with respect to its interaction with the DNA binding site (Figures 4A and B). The atomic analysis of this theoretical complex reveals that atomic interactions between the C terminal regions of the Runt domains and their DNA binding sites are similar to the ones reported in the crystal structure, including Arg80, Arg142, Thr 169, Asp 171, Arg174 and Arg177 [residue numbers as reported in Bravo et al. paper (11); (Figure 4B)]. The model also suggests that the dimer may be maintained by stabilizing interactions between the two molecules of CBFβ (Figure 4C). The proposed structural model predicts favourable interactions between anti-parallel domains of CBFβ that allow optimal electrostatic pairing between polar residues (Figure 4D).

DISCUSSION

It was previously assumed that Runx1 binding to DNA principally involved the interaction of a single Runx1 domain with one isolated TGYGG core sequence (5,6). However, we have made the novel observation that two molecules of Runx1 can assemble together on palindromic sequences containing two closely spaced inverted repeats of a TGYGG core sequence separated by 4 bp. In this arrangement, there is a 2 bp overlap between the 8 bp TGYGGNNN DNA sequences contacted by the individual Runx1 complexes. Runx1 bound most efficiently as a dimeric complex to DNA elements conforming to the consensus TGTGGTTT or TGTGGTCA motifs that represent the most commonly encountered Runx1 binding motifs. Binding of CBF to the Ideal consensus palindromic Runx1 site TGTGGTTAACCACA was more efficient than to the GM450 element, which lacks the preferred TGYGG motifs.

This work is to our knowledge the first study that reports the potential dimerization of CBF complexes on DNA in a conformation similar to that observed for dimers of the Rel homology domains of NFAT and NF-kB family proteins. We are only aware of one other report of Runx1 binding to a palindromic element (42). However, in this instance, only one of the two TGCGG
motifs present within the sequence TGCGGAGACCGCA actually contributed to Runx1 binding (underlined), and there was no evidence for the assembly of dimeric CBF complexes. It has also been reported that the t(8;21) chromosomal translocation product Runx1-Eto is able to bind to direct repeats of Runx1 consensus motifs as found in the consensus sequence TGYGGTTN(0–13)TGC GGT (42). In this situation, the mechanism of dimerization is likely to be different to that showed in our study since in that report it was proposed that oligomerization was mediated by Eto and not by Runx1.

As mentioned above, Runx1 is a distant relative of the Rel and NFAT families of transcription factors, and it shares with Rel domain factors the ability to recognize GG core sequences. Runx1 is also a distant relative of p53 which similarly binds as a dimer to palindromic sequences such as GGGCATGCC (43) which have a conformation resembling both the NF-κB consensus and the dimeric Runx1 site described here. While Runx1 resembles NF-κB and NFAT with respect to its ability to bind simultaneously to both copies of an inverted repeat of the GG motifs, there are significant differences in the nature of the...
intervening sequences that they recognize. Whereas NFAT (18,20,21) and NF-kB (10) each bind as dimers to sequences resembling GGAACATCCC or GAGAGACTCC, Runx1 assembles as a dimer on sequences resembling GGTTAACCC. Runx1 also differs substantially from NF-kB with regard to the fact that NF-kB is not able to use a single Rel domain to bind to a single GG motif. In this respect, Runx1 has more in common with NFAT than with NF-kB. Runx1 and NFAT both frequently cooperate with other classes of transcription factor, but still retain the ability to bind to DNA efficiently to a single GG core element via a single Rel-like DNA-binding domain.

Based on the in vitro binding of purified truncated proteins, there was no evidence for cooperativity in the binding of the recombinant CBF complexes that contain just the Runt domain of Runx1. Nevertheless, it was apparent from the in vivo footprinting that both of the Runx1 sites within the GM450 element were efficiently occupied simultaneously inside activated cells at about half of all sites, raising the possibility of cooperativity in the assembly of the dimeric complex. Furthermore, assembly of dimeric Runx1 complexes appeared to be more efficient with full length naturally occurring assembly of dimeric Runx1 complexes appeared to be more efficient with full length naturally occurring runx1/CBF complexes present in nuclear extracts than with the truncated recombinant runx1/CBFβ. Additional evidence for cooperativity was provided by competition studies showing that a dimeric site was a substantially stronger binding site than an equivalent single binding site. This may be because Runx1 also contains a homodimerization domain at the C terminus that is absent in the recombinant Runx1 (17). It has been previously established that this interaction domain promotes runx1:runx1 interactions between separated runx1 sites. This same domain could also potentially help to stabilize the dimeric complex and support cooperative binding at palindromic runx1 sites. The structure of the C terminal region of Runx1 is unknown, partly because much of it is unstructured, and partly due to technical difficulties associated with preparing full-length recombinant Runx1. However, from the proposed model structure of the dimer it is evident that the C terminal portions of the Runx domains cross over the DNA helices on opposite sides of the DNA. Hence, this allows for the possibility that the CBF dimer encircles the DNA by forming close CBFβ:CBFβ interactions on one side, and Runx1 C terminal interactions on the opposite side of the helix. If the C-terminal region does contain unstructured domains, it is likely to be flexible enough to accomplish this. It is also likely that within the nucleus, the dimeric CBF complex exists as a higher order complex with multiple other regulatory factors that would act to further stabilize such a complex. Further genome-wide analyses will be required to determine how often Runx1 sites within the genome do in fact function as dimeric sites.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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