A complex intronic enhancer regulates expression of the CFTR gene by direct interaction with the promoter

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Abstract

Genes can maintain spatiotemporal expression patterns by long-range interactions between cis-acting elements. The cystic fibrosis transmembrane conductance regulator gene (CFTR) is expressed primarily in epithelial cells. An element located within a DNase I-hypersensitive site (DHS) 10 kb into the first intron was previously shown to augment CFTR promoter activity in a tissue-specific manner. Here, we reveal the mechanism by which this element influences CFTR transcription. We employed a high-resolution method of mapping DHS using tiled microarrays to accurately locate the intron 1 DHS. Transfection of promoter–reporter constructs demonstrated that the element displays classical tissue-specific enhancer properties and can independently recruit factors necessary for transcription initiation. In vitro DNase I footprinting analysis identifies a protected region that corresponds to a conserved, predicted binding site for hepatocyte nuclear factor 1 (HNF1). We demonstrate by electromobility shift assays (EMSA) and chromatin immunoprecipitation (ChIP) that HNF1 binds to this element both in vitro and in vivo. Moreover, using chromosome conformation capture (3C) analysis, we show that this element interacts with the CFTR promoter in CFTR-expressing cells. These data provide the first insight into the three-dimensional (3D) structure of the CFTR locus and confirm the contribution of intronic cis-acting elements to the regulation of CFTR gene expression.

Keywords: CFTR • intronic enhancer • HNF1 • enhancer–promoter interaction

Introduction

Structural and functional analysis of the human genome has revealed that cis-regulatory elements influencing transcription often exist some distance away from relevant basal promoters. These elements may include enhancers, silencers, insulators and locus control regions. Large-scale, functional genomics efforts such as the ENCYclopedia Of DNA Elements (ENCODE) project [1] have begun to annotate the broader context of the human genome by locating and defining these embedded regulatory elements. The cystic fibrosis transmembrane conductance regulator gene (CFTR), which when mutated causes the common genetic disease cystic fibrosis, is associated with a number of potential cis-regulatory sites. These elements often display specific changes in local chromatin structure, and we have previously evaluated the CFTR locus in many cell types for structural features including histone modifications, DNase I hypersensitivity and associated transcription factor binding [2–6]. Here, we use a high-resolution, tiled microarray-based assay, DNase-chip [7], to map DNase I-hypersensitive site (DHS) within 90 kb flanking the CFTR promoter region. We detected a cell-type-specific DHS within the first intron of CFTR that corresponds to a regulatory element that we identified in earlier work [8, 9] and has been confirmed by others [10]. This element (known as 7/8 based on primer sets used to amplify the region [9]) was shown to positively regulate CFTR promoter activity specifically in intestinal cells both in vitro and in vivo. Removal of the element from a human CFTR yeast artificial chromosome (YAC) reduced expression levels of the human gene by about 60% in transgenic mice carrying the YAC but only within the epithelium of the small intestine [8]. Nonetheless, the mechanism of action of this key regulatory element has not yet been explored.

We now utilize this element in transient transfection experiments to show that it functions as a classical, tissue-specific enhancer and can also independently recruit general factors

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necessary for transcription initiation. To determine the nuclear factor(s) interacting with the regulatory sequence, we perform in vitro DNase I footprinting analysis, which reveals a significant protected sequence within which exists a conserved hepatocyte nuclear factor 1 (HNF1) binding site. Expression of HNF1α correlates with CFTR expression, and this transcription factor binds in vitro to another cluster of intrinsic DHS in CFTR [11]. In vivo, HNF1α contributes to the maintenance of normal mouse CFTR expression levels in the small intestine [11]. Here, we show that HNF1 binds to the intron 1 enhancer both in vitro, by electrophoretic mobility shift assay (EMSA), and in vivo, by chromatin immunoprecipitation (ChIP). Moreover, we use chromosome conformation capture (3C) analysis to show that this intronic enhancer interacts with the CFTR promoter in vivo, revealing a three-dimensional structure of the active CFTR locus.

Methods

Cell culture

The human colon carcinoma cell lines Caco2 [12] and HT29 [13], human bronchial epithelial cell line 16HBE14o– [14] and the human embryonic kidney cell line HEK293 [15] were grown in Dulbecco’s Modified Eagle’s Medium (DMEM) (Gibco, Carlsbad, CA, USA) supplemented with 5% fetal bovine serum (FBS). Cells were grown in a humidified atmosphere (37°C) of 5% CO2 and 95% air. The human colon carcinoma cell lines Caco2 [12] and HT29 [13], human bronchial epithelial cell line 16HBE14o– [14] and the human embryonic kidney cell line HEK293 [15] were grown in Dulbecco’s Modified Eagle’s Medium (DMEM) (Gibco, Carlsbad, CA, USA) supplemented with 5% fetal bovine serum (FBS). Cells were grown in a humidified atmosphere (37°C) of 5% CO2 and 95% air.

Primer sequences

All primer sequences and locations used for DNase-chip, RT-PCR, plasmid cloning, mutagenesis, and 3C are listed in the Supporting Information.

DNase-chip

DNase-chip was performed as previously described, with modifications [7]. Briefly, 2–5 × 10⁶ cells were lysed using 0.1% NP-40 buffer. Purified nuclei were exposed to increasing amounts of DNase I (0–30 U; NEB, Beverly, MA, USA) reactions were stopped with 0.1 M ethylenediaminetetraacetic acid (EDTA) and digested chromatin was embedded into InCert agarose plugs (Lonza, Walkersville, MD, USA). Chromatin digestion was determined by pulsed-field gel electrophoresis, and adequately digested samples were sonicated to generate 200–500 bp fragments and the biotinylated ends were captured using 0.25 μg of genomic DNA from the same cell type was labelled with Cy3-dUTP and each was hybridized to ENCODE tiling arrays (NimbleGen, Madison, WI, USA; human genome build 17, May 2004). Hybridization data from three (Caco2 and fibroblasts) or two (16HBE14o– and HT29) experiments were analysed with ACME statistical software [16] using a window size of 500 bp and a threshold of 0.95.

Reverse transcriptase (RT)-PCR

RNA was isolated from Caco2, 16HBE14o–, HT29, HEK293 and primary skin fibroblasts using Trizol (Invitrogen), as per the manufacturer’s instructions. cDNA was generated from 250 ng RNA using the Taqman® Reverse Transcription kit (Applied Biosystems). CFTR expression was assayed using E1 primer set [17]; as a control, β-glucocerebrosidase (GenBank no. M16328) expression was assayed using primers GD67A and GD9B(11).

Plasmid construction

PCR amplification of the CFTR basal promoter and intron 1 putative regulatory elements (referred to as 7/8) was described previously [9]. Reporter constructs for transient expression were generated using the pGL3B vector (Promega, Madison, WI, USA). The CFTR basal promoter was cloned into Nhe I and Bgl II sites of the multiple cloning site, upstream of the luciferase reporter gene. The 7/8 fragment was digested out of the pCRII vector using BamHI, and one or two copies of this fragment were inserted in forward and reverse orientations into the enhancer site of the pGL3B CFTR promoter vector. All subcloned fragments were confirmed by sequencing.

Overlapping fragments of CFTR intron 1 were generated by PCR using cosmids F0424 [6] as a template. All primers included a 5’ Mlu I restriction site, and all amplified products were ligated into the Mlu I site in the multiple cloning site of the promoter-less pGL3B vector upstream of the luciferase reporter gene. All cloned fragments were sequenced to confirm orientation and exclude PCR artefacts.

Mutated plasmid was generated using the QuikChange® Mutagenesis Kit (Stratagene, LaJolla, CA, USA), according to the manufacturer’s protocol.

Transient transfections

Cells were plated 1 day prior to transfection in 96-well plates at a confluence of 5000 cells per well. The cells were transfected with 50 ng of the luciferase reporter DNA and 1 ng of Renilla expression vector (Promega) per well using 0.25 μl/well of lipofectin (Invitrogen), according to the manufacturer’s protocol. The luciferase assay was performed using dual luciferase reagent (DLR) kit (Promega), according to the manufacturer’s protocol. All experiments were corrected for transfection efficiency by normalizing the firefly luciferase reporter gene activity to Renilla luciferase activity. Statistical significance of the results was calculated by unpaired t-tests.

In vitro DNase I footprinting

The intron 1 DHS 7/8 fragment was PCR-amplified using primers TSR7 and TSRS8 and then cloned into pCRII (Invitrogen). Cleavage of this vector with Xho I or Sac I enabled Klenow DNA polymerase fill-in with [α-32P]-dCTP to label the sense or antisense strand, respectively. DNase I footprinting experiments were then performed as described previously [9].
**In vitro transcription/translation of HNF1**

The mouse HNF1α expression plasmid was made by PCR by amplifying the gene from the pBlus-mHNF1α vector kindly donated by Dr. Gerald Crabtree (Stanford University, Stanford, CA, USA). Human HNF1β was cloned using Caco2 cDNA and specific primers. The mHNF1α gene was subcloned into pCRII (Invitrogen) and hHNF1β into pCR-Script (Stratagene); each sequence was verified and cloned into the pcDNA3.1 vector (Invitrogen) using Not I and Hind III (mHNF1α) or Xho I and Not I (hHNF1β).

Wild-type HNF1α and HNF1β proteins were produced using TNT® T7 Quick Coupled Reticulocyte System (Promega), as described in the manufacturer’s protocol. To confirm the presence of translated protein, 5 μl of the reaction volume was resolved on a 10% SDS-PAGE gel and transferred onto nitrocellulose. Immunoblot analysis was performed by probing first with 0.25 μg/ml goat anti-HNF1α primary antibody (Santa Cruz Biotech, Santa Cruz, CA, USA; sc-6547) and anti-goat horseradish peroxidase (ECL) detection reagent (Amersham, Piscataway, NJ, USA). rabbit anti-HRP-conjugated secondary antibody (Sigma, St. Louis, MO, USA). Labelled DNA fragments (~100,000 cpm) were annealed and labelled with [32P]-dCTP. Complementary single-stranded oligonucleotides (see Fig. 4 for sequences) were used. The samples were subsequently diluted to a concentration of 100 ng/ml. A Taqman® probe and reverse primer was designed that was specific to the Hind III fragment encompassing the CFTR promoter. Multiple forward primers were then designed that were specific to different Hind III fragments across the CFTR locus. Using a dilution series of digested/re-ligated bacterial artificial chromosome (BAC) DNA template, each forward primer was demonstrated to function with the ‘fixed’ Taqman® probe and reverse primer to amplify with approximately 100% efficiency. To quantify ligation events within 3C samples, 200 ng of 3C template were used per Taqman® qPCR reaction. The ligation efficiency at each site was corrected for the interaction between two Hind III fragments within the ubiquitously expressed excision repair cross-complementing rodent repair deficiency, complementation group 3 (ERCC3) locus, which has been reported to adopt the same spatial conformation in different tissues [20–22].

**Electrophoretic mobility shift assays**

Complementary single-stranded oligonucleotides (see Fig. 4 for sequences) were annealed and labelled with [α-32P]-dCTP by fill-in reactions with Klenow DNA polymerase, prior to purification with microspin G-25 columns (Amersham Biosciences). Labelled DNA fragments (~100,000 cpm) were incubated for 20 min. with Caco2 nuclear extract (5 μg) or in vitro translated (IVT) proteins (5 μg) in a final reaction volume of 20 μl containing 20 mM HEPES pH 7.5, 100 mM KCl, 10 mM MgCl2, 1 mM EDTA, 12% (v/v) glycerol and 1 μg poly(dI-dC). For competition experiments, the proteins were pre-incubated with unlabelled oligonucleotide duplexes at 100-fold excess molar concentrations, for 20 min. at room temperature before addition of labelled DNA. For supershifts, a similar pre-incubation with nuclear extract or IVT proteins was included with addition of 1 μg of antibodies against HNF1α (Santa Cruz; sc-6547) or HNF1β (Santa Cruz; sc-7411). The samples were resolved on a 4% polyacrylamide gel at 4°C for 2 hrs at 300 V in a 0.5× TBE buffer (1× TBE is 89 mM Tris, 89 mM boric acid and 2 mM EDTA). Following electrophoresis, the gels were dried and exposed to a Phosphormager screen (GE Healthcare, Piscataway, NJ, USA).

**Chromatin immunoprecipitation (ChIP)**

In total, 1 × 10⁷ post-confluent Caco2 cells were trypsinized, resuspended in DMEM and cross-linked with 1% formaldehyde for 10 min. The cross-linking was stopped by the addition of glycine to 0.125 M. The cells were washed with cold PBS and lysed in 1 ml of 1% SDS, 10 mM EDTA, 50 mM Tris/HCl, pH 8.1 and 1× protease inhibitor cocktail (Roche, Indianapolis, IN, USA). Chromatin was sonicated to an average size of 500 bp. Chromatin was sonicated to an average size of 500 bp.

Immunoprecipitations were performed overnight at 4°C using 200 μl chromatin that was diluted 1:10 with ChIP dilution buffer (0.01% SDS, 1.1% Triton X-100, 1.2 mM EDTA, 16.7 mM Tris-HCl, pH 8.1 and 167 mM NaCl) and 10 μg of either rabbit anti-HNF1 antibody (Santa Cruz; sc-8986) or rabbit IgG (Santa Cruz; sc-2027). The complexes were collected using 60 μl Protein A/Salmon sperm agrose beads (Upstate, Billerica, MA, USA), washed several times according to the manufacturer’s protocol and eluted with 1% SDS and 0.1 M NaHCO3. Cross-links were reversed at 65°C overnight, and the samples were treated with RNase (10 μg/ml) and proteinase K (40 μg/ml) before phenol/chloroform extraction and ethanol precipitation. The samples were resuspended in 0.5× TE and enrichment was analysed using Taqman® primer and probe sets and a Fast 7500 Real-Time PCR machine (Applied Biosystems).

**Chromosome conformation capture (3C)**

Chromosome conformation capture (3C) was performed as described previously [18, 19], with minor modifications. Briefly, 1 × 10⁷ cells were fixed with 2% formaldehyde for 10 min. at room temperature. The cells were lysed in 5 ml cold lysis buffer (10 mM Tris pH 8, 10 mM NaCl, 0.2% NP-40 and 1× protease inhibitor cocktail) and the nuclei were collected by centrifugation. Following extraction with 0.3% SDS, chromatin was digested overnight with 2000 U Hind III. Ligations were performed in a total reaction volume of 6.5 ml, using 100 U T4 DNA ligase (Roche), and incubated at 14°C for 4 hrs followed by 30 min. at room temperature. Cross-links were reversed by proteinase K treatment at 65°C overnight. The samples were purified by phenol/chloroform extraction followed by ethanol precipitation and then resuspended in 150 μl H2O. The concentration of each sample was determined by SYBR green quantitative PCR (qPCR), using the B13F/B13R primer set (amplicon found within a Hind III fragment; see Supporting Information), and each sample was compared with a genomic DNA reference of known concentration. The samples were subsequently diluted to a concentration of 100 ng/μl. A Taqman® probe and reverse primer was designed that was specific to the Hind III fragment encompassing the CFTR promoter. Multiple forward primers were then designed that were specific to different Hind III fragments across the CFTR locus. Using a dilution series of digested/re-ligated bacterial artificial chromosome (BAC) DNA template, each forward primer was demonstrated to function with the ‘fixed’ Taqman® probe and reverse primer to amplify with approximately 100% efficiency. To quantify ligation events within 3C samples, 200 ng of 3C template were used per Taqman® qPCR reaction. The ligation efficiency at each site was corrected for the interaction between two Hind III fragments within the ubiquitously expressed excision repair cross-complementing rodent repair deficiency, complementation group 3 (ERCC3) locus, which has been reported to adopt the same spatial conformation in different tissues [20–22].

**Results**

A tissue-specific DHS exists 10 kb into the first intron of **CFTR**

Previous DHS-mapping experiments using restriction enzyme digestion and Southern blotting with labelled **CFTR** probes revealed a prominent DHS approximately 10 kb into the first intron of **CFTR** at 185 +10 kb (185 is the last base of exon 1) and weaker DHS at 185 +12 kb and at +14 kb. The DHS were only apparent in certain cell types, including the human colon carcinoma cell lines Caco2 and HT29, the human pancreatic adenocarcinoma cell line Capan1 and human primary male genital duct cells [9]. Each of these cell types express significant amounts of **CFTR** mRNA. However, in other cell types that express **CFTR**, in particular those
derived from the airway epithelium, such as the human bronchial epithelial cell line 16HBE14o– (Harn’s group, unpublished observations), no significant DHS was evident in this region. To more accurately define these intron 1 DHS, we used DNase-chip, a high-resolution assay for DHS using tiled NimbleGen microarrays, which can define DHS locations much more precisely than is possible by conventional methods. In Caco2 and HT29 cells, a single strong DHS was apparent 10 kb into the first intron (Fig. 1A).

Fig. 1 DNase-chip detects cell-type-specific DHS 10 kb into the first intron of CFTR. (A) Shown are DHS tracks in four different cell types: CFTR– primary human skin fibroblasts, CFTR+ human bronchial epithelial cell line 16HBE14o– and CFTR+ human colon carcinoma cell lines Caco2 and HT29. Significant peaks representing DHS are apparent in the 5′ promoter region of CFTR+ cells, whereas Caco2 and HT29 cells display another significant DHS within the first intron. Peak height is a measurement of –log10 (P-value) between 0 and 16, as determined by ACME (see Methods). The sequence associated with the identified intronic DHS is shown in panel B. Highlighted sequence corresponds to previously identified positive regulatory element 7/8 [9]. Sequence is numbered based on GenBank reference sequence AC000111. (Sequence corresponds to bases 116916908–116918007 of human genome build 17.) (C) CFTR expression in cell types used in this study measured by RT-PCR analysis compared with a β-glucocerebrosidase housekeeping gene control.
This DHS corresponds exactly to the previously defined 185 + 10 kb DHS and overlaps the 7/8 regulatory element [9] previously shown to positively regulate CFTR promoter activity (Fig. 1B). Interestingly, though a strong DHS is evident at the CFTR promoter in Caco2, HT29, and 16HBE14o– cells, the intron 1 DHS is only apparent in the intestinal cells. Moreover, the 185 + 10 kb DHS was the only significant DHS observed within the first intron, indicating that either this is the primary region within the first intron containing cis-regulatory elements or that the cells have evolved in culture and the +12 and +14 kb DHS are now less evident.

The 185 + 10 kb DHS contains a cell-type-specific enhancer of the CFTR promoter

We previously showed that the core of the cis-element within the 185 + 10 kb DHS lies within the 176 bp 7/8 fragment that augments the activity of the basal CFTR promoter in transient transfections of Caco2 cells [9]. These assays showed that this element increased promoter activity both alone and in the context of a larger cloned fragment, demonstrating its function in a position-independent manner. However, we did not determine whether this core had the classical enhancer property of functioning in each orientation. To assay for enhancer activity in transient transfection experiments, we generated a series of pGL3-derived constructs in which a 787 bp CFTR basal promoter fragment drives the expression of the firefly luciferase reporter gene. Each construct has a derivative of the 7/8 regulatory element cloned into the enhancer site of this vector. Single and double copies of 7/8 were cloned in both forward and reverse orientations. To determine tissue-specificity of 7/8 activity, these constructs were transfected into 16HBE14o– bronchial epithelial cells and Caco2 cells that both express CFTR and into human embryonic kidney cells (HEK293) that do not.

These luciferase assays show that one copy of the 7/8 element increases the firefly luciferase activity three-fold over the level of the CFTR promoter alone in Caco2 cells (Fig. 2A). These experiments were repeated in HT29 cells, with similar results (data not shown). This effect is cell-type-specific, as it is not observed in 16HBE14o– cells or HEK293 cells. Equivalent results were seen when the 7/8 fragment was cloned in a reverse orientation, suggesting that it is, indeed, a true enhancer element. Additionally, two copies of the 7/8 enhancer in either orientation show a synergistic effect, increasing CFTR promoter activity by about 10-fold over the single copy element.

The CFTR 7/8 element can recruit factors necessary for transcription initiation

Having confirmed the enhancer function of the 7/8 element, we next evaluated whether regions of the 185 + 10 kb DHS could recruit the necessary factors for transcriptional activation independent of the basal CFTR promoter. We cloned overlapping fragments of intron 1, including the 185 + 10 kb DHS region into the promoter-less pGL3B vector upstream of the firefly luciferase reporter gene. These plasmids were transiently transfected into Caco2 cells and assayed for their ability to activate reporter gene transcription (Fig. 2B). The construct containing one copy of the 7/8 sequence (pGL3B Intr1–7/8) showed strong ‘promoter’ activity, about seven-fold greater than that of the basal CFTR promoter. A construct containing 7/8 with an additional 882 bp of upstream sequence (pGL3B Intr1–7F) showed a similar level of activity. This activity was abolished when this fragment was cloned in a reverse orientation (pGL3B Intr1–7R), suggesting directional function of the 7/8 element. Constructs containing 7/8 with additional downstream sequences did not show ‘promoter’ activity, which may indicate that the inclusion of sequence downstream of 7/8 disrupts luciferase expression in the context of these transient assays, either by recruiting repressive factors or by inhibiting translation of the luciferase transcript.

HNF1 binds an element within the CFTR intron 1 DHS in vitro and in vivo

Previous studies demonstrated that the 185 + 10 kb DHS element binds specific protein complexes in Caco2 cells, though the identity of the factors was not revealed. To identify potential transcription factor-binding sites across the DHS 185 + 10 kb region, in vitro DNase I footprinting was performed (Fig. 3A). Two protected regions were evident upon binding of Caco2 nuclear extracts to a radiolabelled probe spanning the sense strand of 7/8. The protected regions encompass 30 (footprint 1, FP1) and 23 (footprint 2, FP2) nucleotide sequences and suggest a complex pattern of DNA–protein interactions within this site. These data are consistent with footprints identified in our previous analysis of this region, utilizing, in addition to Caco2 cells, nuclear extracts from primary male genital duct epithelial cells [9]. At the time of the original analysis, we were unable to predict the candidate transcription factors involved at this site due to the relative paucity of defined binding sites in available databases. We again performed in silico analysis for putative transcription factor-binding sites using MatInspector (http://www.genomatix.de). Within FP1, a strong predicted binding site for HNF1, which recognizes the consensus sequence GTTAATNATTANC [23], was identified on the antisense strand and it resides within a region of high sequence conservation between a number of species (Fig. 3B), suggesting the presence of a functional element.

To determine the ability of HNF1 to bind at the 185 + 10 kb DHS, EMSAs were carried out using nuclear extracts from Caco2 cells. The nuclear extracts were incubated with an α-32P-labelled double-stranded DNA probe that spans the consensus HNF1-binding sequence, and the mobility of DNA–protein complexes was examined. Two major DNA–protein complexes (A and B) were observed, as indicated by retardation in gel mobility, suggesting that protein–DNA interactions occur at this site (Fig. 4A). These interactions were specific, as the binding of protein complexes...
was effectively competed with a 100-fold molar excess of the corresponding unlabelled probe.

To determine whether HNF1 proteins are present within the complexes bound to DNA, we performed supershift analysis using antibodies directed against HNF1α and HNF1β. Both factors bind to the same consensus DNA sequence and can either homodimerize or heterodimerize [23]. Both HNF1α and HNF1β are produced in Caco2 cells, although the α form is produced at higher levels than the β form (data not shown). The migration of complex A was retarded upon pre-incubation of the nuclear extracts with an antibody specific for HNF1α alone, but not with one specific for HNF1β, suggesting both that HNF1α is a dominant factor present within the Caco2 protein complexes that are formed within the 7/8 enhancer and that other factors are also present (Fig. 4A). The specificity of DNA–protein interactions with HNF1 was confirmed using a probe with a 2 bp mutation within the core of the HNF1-binding site. Pre-incubation of the nuclear extracts with the mutated unlabelled probe competitor had no effect on complex A, suggesting that binding of these factors to DNA requires an intact HNF1-binding motif (Fig. 4A). However, the faster migrating complex B may include other DNA-binding proteins. No clear consensus site was identified in FP2, although *in silico* analysis with the Alibaba 2.1 (http://www.gene-regulation.com) transcription factor search engine, but not with MatInspector.

Fig. 2. Transient transfections of pGL3 reporter vectors reveal enhancer and ‘promoter’ activity. (A) Caco2, 16HBE14o–, and CFTR– human embryonic kidney cell line HEK293 were transfected with constructs with 7/8 cloned into the enhancer site of the vector as a single copy or double copy and either in the forward or in the reverse orientation. A 787 bp fragment corresponding to the CFTR basal promoter used previously [9] was cloned into the promoter site of pGL3B. Renilla luciferase vector was used as a transfection control in all experiments; data shown are relative to the CFTR basal promoter vector without inserted enhancer sequence. (B) Intron 1 fragments cloned into the promoter site of pGL3B were transfected into Caco2 cells. Data shown are relative to CFTR basal promoter vector; error bars represent standard error of the mean (n = 12), *P < 0.01 using unpaired t-tests to analyse the difference from promoter-only vector.
predicted a CCAAT enhancer binding protein (C/EBP)-binding site. EMSA analysis of FP2 confirmed the formation of a DNA–protein complex, yet it was not supershifted with a C/EBP-specific antibody (data not shown). Moreover, no protein complexes were observed when an oligonucleotide probe spanning both FP1 and FP2 with the HNF1 site mutated was radio-labelled and used in EMSA experiments with Caco2 nuclear extracts (not shown), suggesting that destroying the HNF1-binding site eliminates possible binding of any other transcription factors in this region.

To further confirm the identity of the proteins and their direct DNA binding at 7/8, EMSA experiments were performed using IVT HNF1α/H9251 and HNF1β/H9252 proteins (Fig. 4B). In this case, both factors were able to bind and form complexes with the DNA probe. Pre-incubation of IVT proteins with specific antibodies also resulted in the supershift of both HNF1α/H9251- and HNF1β/H9252-containing complexes. This suggests that the recombinant HNF1β protein in addition to HNF1α can bind directly to this DNA element, although not in nuclear extracts where other factors are present.

To demonstrate in vivo binding of HNF1 to this element, ChIP was performed using an antibody that recognizes an epitope present on both forms of HNF1, as the specific antibodies used for EMSA were not suitable for ChIP analysis. Several regions of CFTR were analysed for HNF1 binding in Caco2 cells, including 2 kb upstream of the translation start site, the CFTR basal promoter, +5.7 kb into the first intron, the 185 + 10 kb DHS region and +15 kb into the first intron (Fig. 5). The α1-antitrypsin (AAT) promoter provided a positive control for HNF1 binding [24, 25]. The intronic enhancer element located in the 185 + 10 kb DHS is significantly enriched in HNF1 immunoprecipitation of Caco2 chromatin over an isotype-matched IgG control, establishing in vivo binding of HNF1 to this element. It is also apparent that the basal promoter region of CFTR is enriched at about the same level as the AAT promoter control. Although at low stringency, the AliBaba 2.1 program (http://www.gene-regulation.com) predicts three potential HNF1-binding sites within 1 kb of the CFTR basal promoter, whereas the MatInspector (http://www.genomatrix.de) program does not identify these sites. Moreover, none of these sites bound HNF1 in vitro when evaluated by EMSA (data not shown). Thus, this enrichment could be a result of a direct interaction between the CFTR basal promoter and the distal elements that directly bind HNF1, such as the intronic enhancer at 185 + 10 kb. HNF1 ChIP assays on 16HBE14o- and skin fibroblast chromatin showed no enrichment within the intronic DHS (data not shown).

Destroying the HNF1-binding site abrogates enhancer and ‘promoter’ activity of 7/8

In Fig. 4, we showed that the alteration of two bases in the consensus binding site for HNF1 abolished protein–DNA interactions
in EMSA experiments. Next, we mutated the same two bases within the HNF1-binding site in fragment 7/8 cloned into either the enhancer or the promoter site of the pGL3 reporter vector and assayed them for activity (Fig. 6). Figure 6 shows that destruction of the HNF1 site completely abolishes both the enhancer (Fig. 6A) and the ‘promoter’ (Fig. 6B) activity.

The CFTR intronic enhancer directly interacts with the promoter

As our ChIP analysis suggested that the HNF1-containing complex forming at the 7/8 enhancer might also be interacting with
the CFTR promoter, we performed 3C using a Taqman® reverse primer and probe within a Hind III fragment encompassing the basal CFTR promoter and forward primers on a 5’ fragment and fragments within the first, third and seventh introns (Fig. 7). Using 3C, we measured the relative interaction frequency between the CFTR basal promoter and both 5’ and 3’ distal regions using real-time PCR in Caco2 (intestinal) and 16HBE14o– (airway) cells that express CFTR and in skin fibroblasts that do not. In 16HBE14o– cells and fibroblasts, in which no interaction between the first intron and the promoter is expected (because the intron 1 DHS is absent), the interaction frequency gradually decreases with Hind III fragments that are further from the promoter fragment, as the chance of random interactions decreases. However, in Caco2 cells in which the 7/8 enhancer is active, interaction frequencies between the promoter and the intron 1 fragments are all considerably elevated, and the typical gradual decrease in interaction frequency does not occur, revealing that, in vivo, these fragments are directly interacting with the promoter. The fragment with a Hind III site closest to the observed intron 1 DHS (see DHS bar above Fig. 7, panel A) displayed a significantly higher interaction frequency than a fragment lying equidistant to the basal promoter on the opposite side from intron 1, at approximately 10 kb 5’. This demonstrates that the observed interaction frequency of the first intron is not a result of chance random interactions.

Discussion

Our data suggest a model whereby HNF1 binds to the intronic enhancer at 185 + 10 kb and mediates a loop that functions to augment the recruitment of RNA polymerase II and other general transcription factors to the CFTR promoter, thus increasing CFTR transcription in intestinal epithelial cells. Chromatin loop structures have been shown to play a role in the differential expression of genes at a number of loci and in diverse cellular model systems including erythroid differentiation and T-cell development [26–31]. These loops contain cis-regulatory elements that often exist at large genomic distances from the interacting promoter. Several nuclear factors have been implicated in mediating these higher-order chromatin structures, including factors that are expressed both ubiquitously and in a tissue-specific manner. The expression of CFTR is, with a few exceptions, limited to the chloride-secreting epithelia of various tissues [17, 32–35], yet extensive analysis of the greater CFTR promoter region has not revealed the elements that confer this spatial expression pattern [36–38]. The promoter, which lacks a TATA box, contains potential regulatory elements such as AP-1- and Sp1-binding sites [38], a cAMP response element (CRE) that binds CRE-binding protein (CREB) upon protein kinase A activation [39, 40] and an inverted CCAAT element (Y box) that binds CCAAT enhancer binding protein (C/EBP) [41]. Additionally, NF-κB has been shown to bind about
Fig. 7 The 7/8 enhancer directly interacts with the CFTR promoter. Chromosome conformation analysis of CFTR+ Caco2 (A), CFTR+ 16HBE14o− (B) and CFTR− primary skin fibroblasts (C) show higher interaction of the first intron with the promoter in cells with the intron 1 DHS. Cross-linked chromatin from each cell type was digested with Hind III (sites indicated by dashed lines) and re-ligated. The interaction frequency between a fixed Hind III fragment at the CFTR promoter and Hind III fragments 5′ and within the first, third and seventh introns was measured by Taqman® quantitative PCR. Each reaction was normalized to a control region in the ERCC3 gene, as previously described [31]. The amplification efficiencies of all primer sets were verified using Hind III-digested CFTR and ERCC3 BACs. Each experiment was repeated at least twice; a single representative experiment is shown with PCR reactions performed in triplicate. Error bars represent standard error of the mean (n = 3).
binding to the CFTR intronic enhancer may be playing several
functions: recruiting factors responsible for modifying the chromatin
structure of the enhancer region so that other nuclear factors may
bind, and/or directly recruiting general transcription factors to the
CFTR promoter to increase transcriptional activity. Evidence for
HNF1 binding to the latter role is seen in the ability of the 7/8
enhancer to enable transcription initiation when cloned into the
‘promoter’ site in transient transfection assays. There are, how-
ever, no data to suggest that this intronic element acts as a clas-
sical promoter within the context of chromatin. Thus, we suggest
that the in vitro activity is a specific feature of certain enhancers
that coordinate transcription with endogenous promoters. Indeed,
other known regulatory elements within the CFTR locus also
exhibit this activity in transient assays (data not shown). Several
co-factors have been shown to associate with HNF1α, including
the histone acetyltransferase complex CREB-binding protein
CBP/p300 and the CBP-associated factor P/CAF [49], and these
may also play a role in CFTR promoter activity. In a recent study,
we measured the levels of histone acetylation and methylation at
the CFTR promoter and found enrichment of histone marks asso-
ciated with active chromatin in both Caco2 and primary male gen-
ital duct cells [2]. Our in vitro footprinting analysis of the intron 1
enhancer suggests that other nuclear factors are associated with
this region, yet at this time, we have not identified which factors
are cooperating with HNF1.

Thus, the intron 1 enhancer element is likely the focus for a
complex configuration of multiple factors including HNF1. An
additional regulatory element lying 2 kb 3’ to the site that we char-
acterize here was recently also shown to bind HNF1 in Caco2 cells
[50]. This element likely corresponds to the weak DHS that we
observed previously at 185 + 12 kb [9]. Yet, our current analysis
utilizing tiled microarrays clearly shows that the 185 + 10 kb site
is a much stronger DHS and is thus most likely the site of greatest
regulatory activity in the first intron.

It is also probable that other cis-regulatory elements exist else-
where within the CFTR locus that play a coordinating role in regu-
larizing transcription with the 7/8 enhancer. Our previous analysis of
the CFTR locus revealed a number of putative cis-regulatory ele-
ments, some of which also bind to HNF1 [11]. Hence, it is likely
that, although the 185 + 10 kb intronic enhancer plays a key role
in augmenting the recruitment of general transcription factors and
RNA polymerase II to the promoter in the intestinal epithelium,
other distal elements also physically interact in a complex multi-
ple-loop configuration that includes the CFTR promoter in a trans-
criptionally active chromatin hub. Understanding the mechanism
of action of CFTR enhancer elements and associated nuclear fac-
tors may provide us with novel tools or methods to modulate
CFTR gene expression for therapeutic benefit.

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Supporting Information

Additional Supporting Information may be found in the online ver-
sion of this article.

Table S1 Primers

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