Structural basis for genome wide recognition of 5-bp GC motifs by SMAD transcription factors

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Smad transcription factors activated by TGF-β or by BMP receptors form trimeric complexes with Smad4 to target specific genes for cell fate regulation. The CAGAC motif has been considered as the main binding element for Smad2/3/4, whereas Smad1/5/8 have been thought to preferentially bind GC-rich elements. However, chromatin immunoprecipitation analysis in embryonic stem cells showed extensive binding of Smad2/3/4 to GC-rich cis-regulatory elements. Here, we present the structural basis for specific binding of Smad3 and Smad4 to GC-rich motifs in the goosecoid promoter, a nodal-regulated differentiation gene. The structures revealed a 5-bp consensus sequence GCC(GC)(CG) as the binding site for both TGF-β and BMP-activated Smads and for Smad4. These 5GC motifs are highly represented as clusters in Smad-bound regions genome-wide. Our results provide a basis for understanding the functional adaptability of Smads in different cellular contexts, and their dependence on lineage-determining transcription factors to target specific genes in TGF-β and BMP pathways.
The transforming growth factor β (TGF-β) family of cytokines regulates critical processes during the lifecycle of metazoans, with important roles during embryo development, tissue homeostasis, regeneration, and immune regulation. The main TGF-β signal transduction mechanism involves Smad transcription factors, and mutations in the components of this pathway are responsible for various inherited and somatic diseases. The receptors for TGF-β, nodal, activin, myostatin, and other family members are membrane serine/threonine kinases that phosphorylate and activate Smad2 and Smad3, whereas analogous receptors for the bone morphogenetic proteins (BMPs) preferentially phosphorylate and activate Smads 1, 5, and 8. These receptor-activated Smads (R-Smads) form heterotrimeric complexes with Smad4, which is required for the transcriptional regulation of most target genes. The Smad complexes are recruited to sites throughout the genome by cell lineage-defining transcription factors (LDTFs) that determine the context-dependent nature of TGF-β action. The first identified member of this class of Smad partners is FoxH1 (previously known as FAST1), which binds to nodal-activated Smad2/3–Smad4.

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**Fig. 1** Motif identification in Gsc promoter using ChIP-Seq and CRISPR/Cas9. a) Schematic representation of the proximal promoter and distal enhancer sites of mouse and human Gsc showing the GC-rich region, CAGAC, and the FoxH1 binding sites. b) Heatmap of ChIP-Seq tag densities for Smad2/3 (GSM178291462) and Smad4 (GSM2746361, this work) located at −600 bp from the TSS of Gsc, showing how the signal is centered at the GC-rich region. Coordinates referred to the mm9 genome assembly. Data are displayed for chromosome 12 between 105,711,900 and 105,711,700 bases. c) Scheme of the CRISPR/Cas9-mediated mutagenesis of the Gsc proximal promoter region. CRISPR-mediated deletions of a 10 bp region of the Gsc GC1 site (Clone C1) and of the GC1-FoxH1 region (155 bp, Clone C2) are indicated with dashed lines. The DNA sequence of targeted regions is represented as blue horizontal bars. Deletions were confirmed by deep sequencing and TA cloning (Supplementary Fig. 1e, f). d) The effects of the deletions are shown as the relative mRNA levels of Gsc and Smad7 used as a control of the TGF-beta signaling pathway. qRT-PCR analysis of Gsc mRNA expression in wild type (WT) or Gsc mutant clones and of Smad7 expression in activin A- (green) or SB431542- (gray) treated d3 cells. Gene expression level is normalized to WT samples, n = 3. Error bars represent s.e.m., P < 0.05, Mann-Whitney test using Prism 6 software (GraphPad Software). e) Smad4 ChIP-qPCR data (n = 2) in ES cells showing that the 10 bp deletion in the GC-rich region abolished Smad4 interaction with this region without affecting Smad4 interaction with the Gsc +6 kb distal enhancer element. Error bar represents s.e.m.
complexes to regulate the expression of goosecoid (Gsc) and other mesendoderm differentiation genes in early-stage embryos⁶⁻⁸. The interaction of Smad2/3–Smad4 with FoxH1 in mesendoderm progenitors, and with the LDTFs MyoD in myoblasts, PU.1 in B-cell progenitors, GATA1 in erythroid progenitors, and C/EBPα in myeloid progenitors⁹, 10, represents a paradigm for the versatile, context-dependent regulation of cell stage transitions by TGF-β family factors.

Smad proteins consist of an N-terminal MH1 domain, which binds to DNA, a linker region with phosphorylation sites for network regulatory inputs, and a C-terminal MH2 domain that is phosphorylated by the receptors and binds other Smad proteins, LDTFs, chromatin readers, and transcriptional co-activators and co-repressors⁹, 4. Early insights into the DNA-binding specificity of Smad proteins came from oligonucleotide binding screens, which identified the palindromic duplex 5′-GTCTAGAC-3′ as a high-affinity binding sequence for Smad3 and Smad4¹¹. The X-ray crystal structure of this sequence bound to MH1 domains of Smads 1, 3, 4, and 5 showed that the MH1 domain recognizes the GTCT motif. GTCT or its complementary extended sequence are referred to as the CAGAC Smad binding element (SBE)¹²⁻¹⁵. Subsequently Smad1 and Smad5 were shown to also recognize GC-rich motifs, termed BMP response element (BRE), in certain BMP-responsive genes¹⁶. This apparent dichotomy of SBEs in the TGF-β vs. BMP pathways was surprising, given the extensive sequence and structural similarity of the DNA-binding β-hairpin across Smad MH1 domains. Moreover, DNase I footprinting analysis of the Gsc promoter in vitro¹⁷, 20 and genome-wide ChIP-Seq experiments in live cells²¹⁻²⁴ showed that Smad3 and Smad4 can also bind to GC-rich regions lacking CAGAC sequences.

The mounting evidence that CAGAC may not be the only, or even the strongest Smad-binding element in TGF-β target genes led us to investigate the interaction of Smad3 and Smad4 with GC-rich DNA sequences. Focusing on the human and mouse Gsc promoter, here we identify specific Smad-binding GC-rich motifs and provide the X-ray crystal structures of Smad3 and Smad4.
bound to several of these motifs. Smad1, Smad5, and Smad8 also
bind to these motifs. A high degree of conformational flexibility
of the β-hairpin allows recognition of different 5-bp GC-rich
sequences. Functional assays confirmed that these promoter sites
are required for the Gsc response to nodal in embryonic stem (ES)
cells. The X-ray crystal structure of Trichoplax adhaerens Smad4
MH1 bound to one of these motifs indicates a high conservation
of this interaction in metazoans. Based on these insights we
MH1 bound to one of these motifs indicates a high conservation
of these residues in the absence of DNA. The SAXS scattering curves corresponding to a merged curve generated from data at several protein concentrations (0.5–4.2 mg mL⁻¹) are shown below. χ² values of the different fittings are indicated. b Superimposition of the 10 best structures determined using the solution scattering curves SAXS refined with the NMR chemical shift values obtained in solution. c Identification of residues involved in DNA binding using NMR titrations. Chemical shift differences are shown as bars (green: AGCCGGCTGCC and blue: AGGCCGCGCCGA, respectively) using a 0.2 weighting of ¹⁵N with respect to ²H. The HSQC data displaying the chemical shifts induced upon binding are shown in Supplementary Fig. 3. d Residues affected upon DNA binding are displayed on the surface of the solution structure of Smad4 MH1 domain shown in b. The electrostatic charge distribution was calculated using PyMol.

Results

Nodal-driven Smads bind to GC regions for Gsc activation. We first determined whether nodal-activated Smad2/3 and Smad4 in mouse ES cells interact with the GC-rich region of the Gsc promoter that binds to Smad4 in vitro²⁰ (Fig. 1a, b). We analyzed nine human genome-wide ChIP-Seq data sets⁹, 21–24 that were available in the NCBI GEO database (Supplementary Fig. 1a, b). The proximal promoter (PP) in human Gsc contains one CAGAC motif, which is located between the FoxH1 site and the TSS, and is not conserved in mouse, whereas the PP in mouse Gsc contains

three CAGA repeats, which lack a crucial bp of the CAGAC SBE motif (Supplementary Fig. 1c).

The analysis revealed binding of Smad2/3 and Smad4 to an area of the Gsc PP that encompasses the GC-rich region and the nearby FoxH1 binding site, but excludes the CAGA and CAGAC sites (Fig. 1a, b; Supplementary Fig. 1a, b). Smad binding also occurred in putative distal enhancer (DE) elements that contain CAGAC sequences (Supplementary Fig. 1a, d). These results showed that in ES cells stimulated by TGF-β signals, Smad2/3, and Smad4 bind to a GC-rich region of the PP of Gsc.

To determine the functional relevance of the GC-rich region, we used CRISPR/Cas9-mediated mutagenesis²⁵, ²⁶ to generate focal deletions in the Gsc PP in mouse ES cells. The homozygous deletion of a 155 bp segment including the FoxH1 binding motif and a large portion of the GC-rich region (Supplementary Fig. 1e) abolished the induction of Gsc by activin A (a ligand for nodal receptors) (Fig. 1c, d). Moreover, the homozygous deletion of a 10-bp segment within the GC-rich region (GCGCCGGGCC), which spared both the FoxH1 and CAGA sites, was sufficient to abolish the Gsc response (Fig. 1d). These deletions in the Gsc locus did not alter the response of a separate Smad target gene, the negative feedback regulator Smad7, in the mutant cell clones, indicating that the cells remain capable of activin signal transduction (Fig. 1d).
Additionally, we provide Smad4 ChIP-qPCR data in ES cells showing that the 10 bp deletion in the GC-rich region abolished Smad4 interaction with this region without affecting Smad4 interaction with the DE elements (Fig. 1e).

**Smad4 binding to GC sequences from the Gsc proximal promoter.** In order to characterize the specific GC-rich motifs recognized by Smad proteins, we first designed two 30-bp dsDNA oligonucleotides (GC1 and GC2 segments, Fig. 2a), and measured the interactions by electrophoretic mobility shift assays (EMSA), using recombinant Smad MH1 domains and Cy5-labeled DNAs (Fig. 2b–d). To narrow down the motifs present in these regions, we applied a sliding window approach (6-bp windows sliding by 2-bp steps, within 20-bp duplexes, Supplementary Table 1) covering the GC1 and GC2 segments. The Smad4 MH1 domain-bound dsDNA oligonucleotides containing the GGCGC/G, GGCCGC/G, and GGCTG sequences (underlined in brown, Fig. 2a, b, and Supplementary Table 1) with an affinity in the nanomolar range, as determined by isothermal titration calorimetry (ITC) (Supplementary Fig. 2b). Smad4 bound the GGCGC oligonucleotide with a \( K_d \) of 160.3±0.2 nM. In parallel experiments, Smad4 bound a CAGAC oligonucleotide with \( K_d \) of 270.5±0.1 nM (Supplementary Fig. 2b).

The Smad3 MH1 domain bound the GC1 oligonucleotide with a \( K_d \) of ~100 nM, as estimated from EMSA (Fig. 2b). Smad1 and Smad8 MH1 domains also bind to these motifs efficiently with \( K_d \) values similar to those displayed for a GC-rich BRE and for the
Structural flexibility of the DNA-binding hairpin. To investigate how Smad4 distinguishes between target and non-target GC-rich sequences, we first analyzed the properties of the MH1 domain devoid of DNA, using nuclear magnetic resonance (NMR) spectroscopy in solution. Flexibility and oligomerization properties of the MH1 domain were also analyzed by acquiring \(^{15}\)N NMR backbone relaxation data at two different protein concentrations. The \(^{15}\)N\{\(^1\)H\} steady-state nuclear Overhauser effect (hNOE)\(^{31}\) values are homogeneous all along the sequence except for the \(\beta_2-\beta_3\) pair, which forms the DNA-binding hairpin\(^3\). The \(\beta_2-\beta_3\) hairpin is more flexible than the rest of the protein. (Fig. 3a) The analysis of the T1 and T2 experiments allowed us to calculate a correlation time \(\tau_c\) of 9.5 ns, which is in agreement with a sample of 15 kDa tumbling as a monomer in solution.

These structural properties were confirmed by small-angle X-ray scattering (SAXS) profiles acquired at the European Synchrotron Radiation Facility (ESRF Grenoble, beamline BM29). The data suggest a maximum protein size of 56.5 ± 1.4 Å, \((D_{\text{max}})\) consistent with a globular monomeric particle in solution with a radius of gyration \((R_g)\) of 15.9 ± 1.0 Å. (Fig. 3a) Remarkably, improving the fitting of the experimental data required refining of the X-ray model. The best results were obtained using the backbone chemical shifts converted to dihedral angles and allowing for some flexibility in the extended N-terminus region of the domain and in the \(\beta\)-hairpin involved in DNA binding. A set of the best 10 structures with the lowest \(\chi^2\) to the experimental SAXS profile\(^{12}\) superimposed to the X-ray structure (PDB: 3QSV) is shown in Fig. 3b.

Overall these results indicate that in the absence of DNA, the Smad4 MH1 domain is well structured and behaves as a monomer in solution. The fold is similar to that determined by X-ray crystallography of the Smad4 MH1 complex with DNA\(^{44}\), except that the DNA-binding hairpin presents conformational flexibility in the absence of DNA. We postulate that the flexibility of the hairpin facilitates access to DNA duplexes with slightly distinct topologies (as defined by different A/T or G/C contents), allowing the conserved hairpin to bind different DNA motifs. This hypothesis is supported by the NMR titration experiments performed with two different DNA motifs (Fig. 3c), which induce similar chemical shifts in the same residues located in and around the conserved \(\beta_2-\beta_3\) hairpin (Fig. 3d). Notably, the sequence of this hairpin is highly conserved in all R-Smads and in Smad4 (Supplementary Fig. 2c)\(^{33}\), suggesting that the flexibility observed in Smad4 also occurs in R-Smads.

High-affinity recognition of the GGCGC motif by Smad4. Given that similar residues in the Smad4 MH1 domain interact with different DNAs, we set to identify which nucleotides directly contacted the protein. We applied high-throughput sparse matrix crystallization approaches using a set of 20 different duplexes, with varying DNA lengths and sequences, and obtained X-ray diffraction data of several complexes of Smad4 MH1 domain and these DNAs. The best diffracting crystals were obtained with DNAs of 16 and 18 bp. A complex with the 18 bp dsDNA oligonucleotide containing the GGCGG motif from the GC1 Gsc segment (Fig. 4a) yielded data at 2.05 Å (refer to Methods section and Table 1). The ASU (space group C222\(_1\)) contains only one Smad4 MH1 monomer and a single DNA strand, while the second monomer and the complementary DNA strand in the biological assembly belong to the neighboring ASU (Fig. 4b).

In the complex, the concave face of the DNA binding hairpin dives into the concave major groove of the duplex DNA containing five base pairs (6-GGCCG-10/9'-GGGCCG-13'). The main DNA-binding region of the protein comprises the loop following the \(\beta_1\) strand, and the \(\beta_2-\beta_3\) hairpin (residues 75–89 in Smad4). The Smad4 \(\beta_2-\beta_3\) hairpin contains three residues Arg81, Glu83, and Lys88, which are strictly conserved in all R-Smad and Smad4 proteins and define specific interactions with three nucleotides in the canonical GTCT motif previously investigated\(^{14, 34}\). In the complex with the 6-GGCCGG-11 site, these three residues also participate in a network of hydrogen bonds with the first four consecutive base pairs of the GGCGC motif.

The electron densities for these functionally important residues, the rest of the protein and DNA are well defined (Fig. 4c). Specifically, the guanidine group of Arg81 side chain coordinates tightly the G6 nucleotide. The orientation of Arg81 is further stabilized by the interaction of Asp79 side chain with the guanidine side of Arg81. In addition, Lys88 side chain is also well defined through interactions with both G7 and G11\(_l\), the latter being located in the complementary chain. The Glu83 side chain carboxyl interacts with C10\(_l\), while the amide part is in the proximity of the C9\(_l\) phosphate. The complex is further reinforced by a set of hydrogen bond interactions between the

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**Fig. 4** Smad4 and Smad3 MH1 domains bound to the GGCGC site. a Schematic representation of the GC1 site with the crystallized regions highlighted in gray. DNA sequences are shown in Supplementary Table 1. b Biological assembly of huSmad4 MH1–GGCGG complex (2.05 Å resolution). The hairpin binding site is circled. The bound Zn\(^{2+}\) and coordinating residues are shown. c A close view of the GGCGC recognition. Distances are shown in Å. The electron density corresponding to the binding region is contoured at 1σ level (2F\(_{o}-F\)). The stereoview representation of the complexes is shown as Supplementary Fig. 9. d Intermolecular contacts. Solid lines indicate hydrogen bonds between protein residues and DNA bases. Dashed lines indicate hydrogen bonds between residues and DNA phosphates. Bases are colored and labeled. e ASU of Trichoplax adhaerens Smad4 MH1 (shown in blue) with the GGCGG site (2.43 Å). f Specific intermolecular contacts of huSmad4 MH1 with the GGCGC site. Distances shown in Å. g Superposition of the Human (green) and Trichoplax (blue and grey) Smad4 MH1 complexes with the same GGCGC DNA. Protein loops displaying minor structural differences are indicated (numbers correspond to the human sequence). h ASU of the huSmad3 MH1 complex with the same GGCGG (2.05 Å). i Expanded view of the GGCGG site with bound Smad3. The electron density corresponding to the binding region is contoured at 1σ level (2F\(_{o}-F\)). j Intermolecular contacts for the huSmad3 MH1 complex with the GGCGG site. k Summary of specific DNA–protein interactions mediated by water molecules for the huSmad4–GGCGC complex. Hydrogen bonds are represented by black dashed lines. Water molecules common to the huSmad3 GTCT complex (PDB:3O2J) are shown in bold green, water molecules present in both Smad4 and Smad3 complexes are shown in green. Analyzed waters were selected as described in the Methods section (numbers correspond to those in the PDB files). l Summary of specific DNA–protein interactions mediated by water molecules for the Smad4–GGCGG complex, as in k. Waters only present in the Smad3 complex are shown in blue. m Superposition of the huSmad3 (tan) and huSmad4 (chartreuse) GGCGG complexes. Minor differences are detected at the a1–a2 and after a3. The DNA-binding site is nearly identical
backbones of Leu78, Gln83, and Ala85, with G11, G9', and C10', covering the 6-GGCGCG-11 area (Fig. 4d) and by a network of well-ordered water molecules, which are tightly bound at the interface of the protein–DNA-binding site (Fig. 4k). Ten of the water molecules are closely coordinated by residues of the β2–β3 hairpin and six bases of the GGGCGCGC region. Additionally, five more water molecules strengthen the interactions of the N-terminal helix α2 with the phosphates of the DNA backbone.

Table 1 X-ray data collection and refinement statistics

|                          | SMAD4-GGCCG 5MEY | SMAD4-GGCT 5MEZ | T_SMAD4-GGCCG 5NM9 |
|--------------------------|------------------|-----------------|--------------------|
| **Data collection**      |                  |                 |                    |
| Space group              | C222             | P21;2;2         | P21;2;2            |
| Cell dimensions          |                  |                 |                    |
| a, b, c (Å)              | 64.17, 79.00, 90.10 | 64.79, 79.06, 114.16 | 37.15, 76.98, 145.04 |
| α, β, γ (°)              | 90, 90, 90       | 90, 90, 90      | 90, 90, 90         |
| Resolution (Å)           | 50.00–2.05 (2.06–2.05) | 30.00–2.98 (2.99–2.98) | 30.00–2.43 (2.44–2.43) |
| R_mer (%)                | 6.7(118.3)*      | 9.4 (91.5)*     | 8.1(182.6)*        |
| I/σI (%)                 | 17.2(1.59)*      | 12.86 (1.68)*   | 12.32(0.80)*       |
| Completeness (%)         | 99.9 (100)*      | 99.7 (100)*     | 98.7(100)*         |
| Redundancy (%)           | 6.2 (6.0)*       | 3.83 (4.06)*    | 4.4 (4.8)*         |
| **Refinement**           |                  |                 |                    |
| Resolution (Å)           | 29.70–2.05       | 29.05–2.98      | 29.43–2.43         |
| No. reflections          | 14,709           | 12,239          | 16,190             |
| R_work/R_free (%)        | 0.222/0.238      | 0.217/0.252     | 0.221/0.251        |
| No. atoms                | 1470             | 2534            | 2656               |
| Protein                  | 979              | 1927            | 1897               |
| RNA                      | 369              | 618             | 738                |
| Zinc ions                | 1                | 2               | 2                  |
| Calcium ions             | 5                | 0               | 0                  |
| Water                    | 92               | 7               | 19                 |
| B-factors                |                  |                 |                    |
| Protein                  | 48.35            | 75.36           | 73.32              |
| DNA                      | 50.76            | 96.29           | 105.01             |
| Zinc ions                | 45.74            | 66.79           | 58.49              |
| Calcium ions             | 107.54           | 0               | 0                  |
| Water                    | 51.75            | 63.97           | 61.92              |
| R.m.s. deviations        |                  |                 |                    |
| Bond lengths (Å)         | 0.009            | 0.010           | 0.010              |
| Bond angles (°)          | 1.00             | 1.05            | 1.02               |

|                          | SMAD3-GGCCG 5OD6 | SMAD3-GGCT 5ODG | SMAD4-GGCCG 5MF0 |
|--------------------------|------------------|-----------------|------------------|
| **Data collection**      |                  |                 |                  |
| Space group              | I4_1             | I4_1            | P4_1             |
| Cell dimensions          |                  |                 |                  |
| a, b, c (Å)              | 105.20, 105.20, 73.24 | 104.99, 104.99, 72.49 | 101.52, 101.52, 45.78 |
| α, β, γ (°)              | 90, 90, 90       | 90, 90, 90      | 90, 90, 90       |
| Resolution (Å)           | 30.00–2.00 (2.05–2.00) | 30.00–2.12 (2.13–2.12) | 30.00–3.03       |
| R_mer (%)                | 9.0 (117.9)*     | 4.9 (66.30)*    | 11.6(98.7)*       |
| I/σI (%)                 | 7.51 (1.20)*     | 14.84 (1.70)*   | 12.06(1.76)*      |
| Completeness (%)         | 99.5 (98.8)*     | 99.7 (98.4)*    | 99.8(98.8)*       |
| Redundancy (%)           | 2.90 (2.95)*     | 3.77 (3.64)*    | 4.9(5.1)*         |
| **Refinement**           |                  |                 |                  |
| Resolution (Å)           | 27.11–2.00       | 29.83–2.12      | 28.25–3.03        |
| No. reflections          | 26,771           | 22,057          | 9298             |
| R_work/R_free (%)        | 0.195/0.235      | 0.200/0.243     | 0.231/0.270       |
| No. atoms                | 2863             | 2647            | 2608             |
| Protein                  | 2034             | 2026            | 1940             |
| RNA                      | 656              | 533             | 656              |
| Zinc ions                | 5                | 2               | 2                |
| Calcium ions             | 0                | 0               | 0                |
| Water                    | 167              | 81              | 4                |
| B-factors                |                  |                 |                  |
| Protein                  | 55.01            | 62.2            | 62.52            |
| RNA                      | 81.93            | 84.12           | 104.88           |
| Zinc ions                | 48.16            | 53.21           | 51.85            |
| Calcium ions             | 0                | 0               | 0                |
| Water                    | 57.41            | 58.73           | 40.44            |
| R.m.s. deviations        |                  |                 |                  |
| Bond lengths (Å)         | 0.01             | 0.01            | 0.008            |
| Bond angles (°)          | 1.00             | 1.06            | 0.97             |

* Values in parentheses are for highest-resolution shell.
These results indicate that the complex interface is highly complementary and that one MH1 protein covers a DNA-binding site of six base pairs. When compared to the structure of the Smad4–GTCT complex, the proteins are nearly identical with minor differences detected in one of the loops and at the first helix (Cα RMSD of 0.39 Å for 113 aligned residues). Most of the differences concentrate at the DNA-binding interface.

We also solved the X-ray crystal structure of Smad4 MH1 in complex with the GGCGG sequence, which is present in both the GC1 and GC2 segments (Fig. 2e). The complex with GGCGG (data at 3.03 Å, Space group P4₃) is similar to that with GGCCG, (Supplementary Fig. 4). The DNA sequence contains the GGCCG motif twice, and each of two bound MH1 monomers interacts slightly differently with this motif, with both binding sites in the biological unit displaying specific hydrogen bonds to either 5- or 4-nucleotides, respectively. Data collection and statistics are shown in Table 1. These additional structures showed that Smad4 can recognize slightly different motifs (GGCCG and GGCGG) in the GC-rich region of Gsc.

**Smad4 recognition of the GGCGG motif across metazoans.** We examined the recognition of the GGCGG site by the Smad4 MH1 domain of the Placozoa Trichoplax, metazoans whose bodies are composed by only four basic cell types and with one of the most divergent among the known Smad4 sequences relative to human Smad4 (Supplementary Fig. 2c). The crystal structure of the Trichoplax Smad4 MH1 domain bound to the GGCGG oligonucleotide reveals that this Smad4–DNA interaction is strictly conserved (Fig. 4c, f) (diffraction data obtained at 2.43 Å, space group P2₁2₁2₁, data collection, and statistics shown in Table 1). A superposition of the human Smad4 and Trichoplax Smad4 complexes shows that both complexes are very similar, and in particular the protein binding mode (Cα RMSD of 0.46 Å for 115 aligned residues). The few observed differences are concentrated in and around two loops that also contain most of the amino-acid sequence differences (Fig. 4g). Collectively, the results show that Smad4 proteins from highly divergent metazoan species interact with the GGCGG motif with similar binding affinity and similar structural contacts.

**Mode of Smad3 binding to the GGCGG motif.** Next, we investigated the binding mode of the Smad3 MH1 domain with the GGCGG motif. Crystals of human Smad3 MH1 domain bound to a 16mer dsDNA oligonucleotide containing the GGCGGC motif yielded data at 2.05 Å resolution using synchrotron radiation (I₄₁₃ symmetry ID29, ESRF, Grenoble), which together with the Smad4 bound to the same site, are the highest-resolution structures determined for a DNA-bound Smad MH1 complex.
domain to date. The asymmetric unit (ASU) of the complex contains two Smad3 monomers and one dsDNA (Fig. 4h). Phases were obtained by molecular replacement using the crystal structure of the human Smad3 in complex with the GTCT motif, (PDB accession code: 1OZJ34). The electron density map obtained after refinement showed residues 10–135 of Smad3, and the dsDNA 16mer (data collection and statistics are shown in Table 1), including the presence of a bound Zn2+, coordinated by His126, Cys64, Cys109, and Cys121.

As in all other MH1 complexes, the three residues strictly conserved (Arg74 and Gln76 located in β2 and Lys81 in β3) are well defined in the structure and participate in a network of hydrogen bonds with the 5-GGGCGG-10 motif. The guandine group of Arg74 is well ordered due to its tight coordination by the backbone and the DNA phosphates. In addition, interactions occurred between Leu71, Gln76, Ser78, and Lys81 to the DNA via bound water molecules, further stabilizing the complex. In the past, it has been suggested that binding of the Smad3 MH1 domain to GC-containing motifs might be unspecific through contacts made from lysine and arginine residues present in helix α2 with the DNA34. However, among the many hydrogen bonds detected from the protein to DNA, only two of them involved two residues located in this helix and were facilitated via water molecules. In fact, Smad3 and Smad4 GGCGC complexes are structurally very similar with a Cβ RMSD of 0.44 Å for 105 aligned residues and both complexes interact directly with the DNA using the β2–β3 hairpin (Fig. 4m).

**Smad3 and Smad4 interactions with the GGCT motif.** The GC-rich region of Gsc contains a conserved GGGTG sequence, which resembles the GTCT-AGAC SBE (Fig. 5a). This GGGTG motif bound Smad3 and Smad4 MH1 domains with high affinity in the EMSA experiments. We determined how the T to G change modified the recognition of the motif by Smad3 and Smad4 using a palindromic version of the motif similar to the canonical SBE. The X-ray crystal structure of the Smad4 MH1 domain in complex with a GGCT-AGAC oligonucleotide was determined at 2.98 Å resolution (space group P212121). The ASU contains two Smad3 complexes, one of them containing the protein in a conformation similar to the canonical SBE. The main interactions with the GTCT motif, similar to what we observed in the GGCGC complex (both conformations are shown in Fig. 5c, d).

The Smad3 MH1 domain complex with the GGCT oligonucleotide was refined at 2.12 Å resolution (space group I4 1) (Fig. 5e). Most of the hydrogen bonds with the GGCT site are very similar to those observed with Smad4 (Fig. 5f, g). The main

![Diagram of Smad4 complexes](image_url)

**Fig. 6 Consensus binding site for GC motifs and DNA shape comparison.** a) Superposition of the human Smad4 complexes with the GGCGC site (shown in green) and the GGCT site (in violet) to the GTCT structure (sky blue) previously determined (PDB: 3QSV). b) Superposition of Smad4 (Top) and Smad3 (Bottom) binding sites. Nucleotides that interact with the protein are highlighted. The Smad4 complexes are colored as in a. Smad3 complexes are displayed in tan (GGCGC site), brown (GGCT), and light blue GTCT (structure previously determined PDB: 1OZJ). c) Structural alignment of the DNA-binding sites observed for the six complexes, with the contacts to the three conserved residues R, K, and Q highlighted. d) DNA shape comparison for different complexes. Major groove width (top) and depth (bottom) were calculated using Curves+ for Smad3 MH1–GGCGC complex (tan), Smad3 MH1–GGCT (brown), Smad4 MH1–GGCGC complex (green), and Smad4 MH1 (light blue) bound to the GTCT site (PDB: 3QSV). The analyzed sequences are indicated.
difference is that in the Smad3 complex no hydrogen bonds are observed from the \( \varepsilon \)-amino group of Lys81 to the G6 nucleotide. In this complex, we also detected several tightly bound water molecules at the protein–DNA interface that contribute to the stabilization of the interactions (Fig. 5g).

**Comparative analysis of Smad binding to different motifs.** We analyzed the superposition of the Smad4 MH1 domain bound to different DNAs (Fig. 6a) with \( \alpha \) root mean square deviations (RMSD) of 0.47 Å and of 0.45 Å for the GTCT and GGCT, respectively, for 114 aligned residues. Similar RMSD values were obtained when we compared the Smad3 complexes. Both GGCGC and GTCT structural comparisons reveal differences in the DNA-binding mode, including the number of direct hydrogen bonds to the bases, and in the topology of the DNA structures, (Fig. 6b), although the same protein residues are involved in the interactions (Fig. 6c). The complexes of Smad4 or Smad3 MH1 domain with the ds-GGCCG motif show a narrower but deeper major groove and a wider and less pronounced minor groove (average width of 10.0 and 6.5 Å, respectively for the Smad4 complex) compared to this domain bound to GTCT/AGAC (3QSV) (10.6 and 5.9 Å, respectively). The conformation of the Smad-bound GGCT motif is intermediate between that of GTCT and that of GGCGC. Similar values were measured for Smad3 complexes. The similarities and differences of both topologies were quantified using Curves+33 (Fig. 6d). Similar topologies of GC-rich DNAs have been previously reported in other cases including the binding of several consecutive Zn-finger domains to GC motifs35. These differences in the DNA shape and flexibility provide a good match between the convex face of the DNA-binding Smad hairpin and the concave and narrow major groove of the GGCGC site, allowing the side chains of Arg81 and Lys88 to establish a well-defined set of hydrogen bonds with G nucleotides.

| Motif      | Average number of motifs per 200bp | Percent of ChIP-Seq peaks containing 0, 1, 2, 3+ motifs |
|------------|-----------------------------------|------------------------------------------------------|
| GGCGC/G    | 66%                               | 0, 1, 2, 3+                                           |
| GGCCG      | 39%                               |                                                      |
| GGCTG      | 55%                               |                                                      |
| DAGAC      | 63%                               |                                                      |
| Baseline   | 6%                                 |                                                      |

**Fig. 7** 5GC motifs are enriched in nodal/TGF-β and in BMP-stimulated cells CREs. a Bar representation showing the average number of GC (GGCGC/G, GGCTG, and GGCCG), CAGAC, and DAGAC motifs found in 200 bp-regions of Smad1, Smad3, and Smad4 ChIP-Seq data up to 1000 bp from the TSS (shown in brown, beige, and green, respectively) or from the baseline data set (gray). AGAC motifs preceded by nucleotides different from C are included under the DAGAC name. b Stacked bar plots showing the number of motifs per 200 bp region (as %) found in Smad1, Smad3, in Smad4-bound ChIP-Seq data, considering the promoter regions as above. Smad4 data corresponding to HepG2 cell lines are shown for comparison of the distribution of motifs to ES. Regions are classified as 0, 1, 2, 3, or more sites. Left: GGCGC/G, GGCCG, GGCTG sites together vs. baseline, colored in blue. Right: CAGAC motifs vs. baseline, colored in brown. c Model of the GC region with the three Smad DNA structures determined. Although based on the sequence, there is an additional potential binding site in this region, only three binding sites are available. Once the first domain is bound (in one of the places), there are only two more sites accessible. With this arrangement, there is no direct contacts between the MH1 domains.

![DNA diagram](image-url)
A consensus GC-rich Smad binding element. The six MH1–DNA complex structures reveal common features of how Smad3 and Smad4 recognize GC motifs in a specific manner. They also illustrate why not all GC motifs can be Smad binding targets. Efficient interactions with GC sites occur only if a G nucleotide is located deep in the major groove, and establishes hydrogen bonds with the guanidinium group of Arg81. This interaction facilitates a complementary surface contact between the Smad DNA-binding hairpin and the major groove of the DNA. The ε-amino group of Lys88 also displays hydrogen bonds with two nucleotides in positions +1 and +2 (also guanines) with respect to the first G. Due to the extended side chain of the lysine, the guanines can be located at either side of the DNA duplex. These conserved sets of contacts define the first three nucleotides in the preferred motifs. Gln83 is more versatile and can form hydrogen bonds using its bi-functional side chain. Indeed, Gln83 interacts with GC in GGGCGC (or CG in GGCGCG) and with an adenosine in GTCT and GGCT.

Additional contacts are observed with the DNA backbone to define an area of interactions with up to 6 bp. The pattern of contacts and the binding mode elucidated here for Smad3 and Smad4 with the GGGCGC motif are slightly different to that recently described for Smad5 bound to the palindromic GGCGCC site.15 In the Smad5 complex, perhaps favored by recent descriptions for Smad5 bound to the palindromic GGCGCC sequence, the first three nucleotides in the preferred motifs. Gln83 is more versatile and can form hydrogen bonds using its bi-functional side chain. Indeed, Gln83 interacts with GC in GGGCGC (or CG in GGCGCG) and with an adenosine in GTCT and GGCT.

The Smad MH1 domain is highly conserved in distantly related metazoa, with no amino-acid sequence differences in the DNA-binding β-hairpin (with the exception of Drosophila Smad2). The structures determined here reveal that the β-hairpin is flexible and adaptable to bind with high-affinity several variants of the consensus sequence GGC(GC)(CG), which we refer to as 5GC SBE. The structures also define the contacts of the Smad3 and Smad4 MH1 domains with specific DNA bases and phosphate groups.

Given the sequence identity of the β-hairpin and the overall high conservation of the MH1 domain in all R-Smad proteins and Smad4, it is not surprising that the various 5GC SBE motifs similarly bind to Smads 2 and 3, which operate in the TGF-β/nodal signaling, Smads 1, 5, and 8, which operate in BMP signaling, and Smad4, which is shared in all of these pathways. Binding of Smads 3 and 4 to the 5GC motifs is of similar or higher affinity than the binding of these proteins to the CAGAC motif GTTG (and to its variant GGCT), which has long been considered as a TGF-β responsive SBE. Smads 3 and 4 also bind to the motif GGGCGC, which is similar to the palindromic GGGGCC sequence previously identified as a BRE for BMP-activated Smad1.

The present results challenge long-standing perceptions that a dichotomy exists between the intrinsic preference of different Smad proteins for different SBEs. The concept that Smad2/3 and Smad4 primarily bind CAGAC SBEs, whereas Smads 1, 5, and 8 prefer GC-rich motifs seem too restrictive in light of the evidence. We propose that all R-Smad and Smad4 proteins recognize G(T)/G/CTG motifs of the classical CAGAC SBE as well as GGC(GC)(CG) motifs of the 5GC SBE. Analysis of ChIP-Seq data sets corresponding to CREs indicates that Smad responsive regions frequently contain both 5GC and CAGAC SBEs, with a higher abundance of 5GC SBEs, which frequently occur as clusters. Additional biochemical and functional studies would need to be done in the future to validate the genome-wide findings and to narrow down the specific regions recognized by Smad proteins case by case.

Smad proteins do not bind to just any 5-bp GC-rich motif, and their structures of Smad–DNA complexes illuminate the basis for the selectivity of these interactions. These insights also suggest how three MH1 domains in a trimeric Smad complex may bind excluding CAGAC motifs). Similar values of cluster enrichment were observed in Smad3 (39% of regions with three or more 5GC motifs vs. 4% of CAGAC), and Smad4 (55% 5GC vs. 6% CAGAC). We also analyzed the 5GC clustering in a Smad4 ChIP-Seq in human hepatocarcinoma (HepG2) cells (Encode ENCF484VVM data set). In this data set, we found similar results with respect to the hESC cluster enrichment, with 63% of the peaks showing three or more 5GC motifs vs. 2% of CAGAC clusters. Finally, a quarter of the promoter regions studied in the three Smad ChIP-seq data sets contain clusters of three or more 5GC SBEs accompanied with CAGAC motifs. Some examples of the identified clusters in TGF-β/nodal and BMP target genes are shown in Supplementary Figs. 6 and 7, respectively. In addition, to explore the 5GC cluster enrichment outside of promoter regions, we also performed these analyses using all Smad4 ChIP-Seq peaks genome-wide (Supplementary Fig. 8).

As revealed by the structures with the 5GC motifs, the fact that specific contacts in the complexes involve nucleotides present in both DNA strands provides an additional plasticity to the interactions but also adds complexity for the identification of these motifs in ChIP-Seq data analysis genome wide, explaining why the GC motifs have remained elusive in these analyses.
to DNA at once. The clustering of multiple SBEs in gene promoter regions may facilitate the cooperative binding of three Smad molecules in the same complex to a target promoter. This is illustrated by a model of three Smad MH1 domains binding to 5GC SBEs in the Gsc PP (Fig. 7c).

The regulation of distinct gene sets by TGF-β and BMP signals is dictated by the differential binding of Smad2/3 and Smad1/5 to master lineage-determining transcription factors, such as FoxH1 in the context of mesendoderm progenitors, and other transcription factors in other cellular contexts.1 LDTFs bound to cognate sites throughout the genome recruit activated Smad complexes to these loci, thus directing where Smads may bind to DNA if appropriate motifs are regionally available in these loci. In this model, the target gene specificity of TGF-β vs. BMP-activated Smads is primarily dictated by the selective affinity of certain LDTFs for Smad2/3 or Smad1/5. The flexibility of the DNA-binding β-hairpin to recognize different variants of the 5GC and CAG SBEs, as shown here, combined with the occurrence of these SBEs in clusters, facilitates the binding of the LDTF-recruited Smad complex to DNA for the assembly of functional transcriptional complexes. This capacity also endows Smad transcriptional complexes with a level of adaptability that may have contributed to establish TGF-β signaling as one of the most versatile and highly conserved signaling pathways in metazoans. Our findings suggest a molecular basis for the functional versatility of Smad complexes in different biological contexts.

Methods

Protein expression and purification. The plasmid (Addgene #14595) encoding the human Smad4 protein was used as the template for cloning different constructs of the MH1 domain to optimize the yield and homogeneity of the recombinant protein. The best results were obtained with the Pro10-Gly140 fragment of Smad4, which was expressed as a N-terminal His-tagged fusion protein followed by a TEV protease cleavage site. The MH1 domain is very stable in solution, showing a melting temperature of 66.2 ± 0.1 °C determined by differential scanning calorimetry and well folded according to the NMR data. The 1D and 2D NMR data of this recombinant domain show a well-dispersed pattern of chemical shifts indicative of a folded sample (1H, 13N-Heteronuclear 2D Single Quantum Coherence, and Transverse Relaxation-Optimized Spectroscopy, HSQC-TROSY).

Two divergent Smad4 MH1 domains, (Ampthalam queenslandica, (Porifera) NCBI Reference Sequence: XP_003388571, Thr52-Thr192; and T. adhaerens (Placozoa), GenBank: EDV21247, Met-1Ser151) as well as the Hu-Smad3 (NCBI Reference Sequence: NP_005893, isoform 1, Met-Pro141), the Smad8/9 domain (UniProtKB: Q99717-1, Thr11-Arg 143), Smad2 (Q99717-1, Thr11-Arg 143) were cloned using a synthesized DNA template with reference values, as well as the 15N edited-NOEY data, corroborated the presence of bound Zn2+ and of four helices and six strands, characteristic of the MH1 fold. These strands are ordered as three anti-parallel pairs: f1βββ, f2βββ, and f3βββ. The presence of many long-range interactions confirmed that, the structure of the MH1 domain is well defined in solution, in the absence of DNA. Chemical shifts have been deposited in the BMRB (entry 26945). For the titration experiments, NUS samples (500 μM for backbone and 15N-NOEY experiments and 100 μM for titrations) were equilibrated in a buffer containing 20 mM sodium phosphate and 100 mM NaCl. All samples were supplemented with 10% D2O and pH adjusted to 6.3. Spectra were acquired using 15N-labeled protein samples at the indicated concentrations with progressively increasing amounts of the unlabeled protein. Chemical shifts perturbation experiments were performed by monitoring of 15N with respect to 2H. NMRPipe 60 and MnmDNR 59 were used for spectroscopy and spectra assignment and analysis was performed with CARA 41.

Chromatin immunoprecipitation. mESCs E14Tg2a.IV (RRID: MMRRC_015890-UCD) were maintained on gelatin (0.1%, Millipore, E008-06)‐coated plates with LIF-supplemented medium at 37 °C with 5% CO2. Basic ES cell medium included 800 U/ml penicillin, 100 μg/ml streptomycin (Invitrogen), 100 μg/ml bovine serum (HyClone, SH30071), 50 U Penicillin and 50 μg/ml Streptomycin (Cellgro, 30-001-CI), 1% non-essential amino acids (Life Technologies, 11140-050), 1% l-glutamine (Life Technologies, 25030081), 100 μM x-mercaptoethanol (Sigma-Aldrich, M5250), 1000 μM l-mlf (Gemini Bio-technology). About 2% pre-cleared chromatin prior to primary antibody addition was kept as input DNA. Magnetic beads were washed, chromatin was eluted, and reverse crosslinked ChIP DNA was dissolved in 10 mM Tris pH 8.0 buffer for further analysis.

ChIP-Seq DNA samples were sequenced on the HiSeq2500 platform. Sequencing reads in FASTQ format were quality assessed by Ribogreen and FastQC v0.11.3 for sequencing integrity and completeness. Additional information is described in Table 1. Cloned products were first screened through aberrant melting temperature of qPCR products, then verified by PCR, TA cloning, Sanger sequencing, and CRISPRe-Plex analysis individually. The mESCs used for this experiment are diploid.

NMR chemical shift assignment and perturbation experiments. NMR data were recorded at 298 K on a Bruker Avance III 600-MHz spectrometer equipped with a quadrupole (1H, 13C, 15N, 2H) resonance cyrogene probe head and a 2-pulse field gradient unit at 298 K. Backbone 1H, 13C, and 15N resonance assignments were obtained by analyzing the TROSY versions of 3D HNCA/CB and HN(CO)CACB experiment pair using (H2O, 1H, 13C, 15N) samples 38. To optimize the quality of the triple resonance backbone spectra, all experiments were acquired as Band-Selective Excitation Short-Transient-type experiments (BEST) with TROSY and non-uniform sampling (NUS), using two different buffer conditions (pH 7.5 and 5.5) to minimize the number of overlapped amides.1 This strategy allowed us to unambiguously assign 116 of the 125 possible amides (131 residues, 6 of them prolines). The comparison of the Smad MH1 (Cz and Cβ) chemical shifts to reference values, as well as the 15N edited-NOEY data, corroborated the presence of bound Zn2+ and of four helices and six strands, characteristic of the MH1 fold. The strands are ordered as three anti-parallel pairs: f1βββ, f2βββ, and f3βββ. The presence of many long-range interactions confirmed that, the structure of the MH1 domain is well defined in solution, in the absence of DNA. Chemical shifts have been deposited in the BMRB (entry 26945). For the titration experiments, NUS samples (500 μM for backbone and 15N-NOEY experiments and 100 μM for titrations) were equilibrated in a buffer containing 20 mM sodium phosphate and 100 mM NaCl. All samples were supplemented with 10% D2O and pH adjusted to 6.3. Spectra were acquired using 15N-labeled protein samples at the indicated concentrations with progressively increasing amounts of the unlabeled protein. Chemical shifts perturbation experiments were performed by monitoring of 15N with respect to 2H. NMRPipe and MnmDNR were used for spectroscopy and spectra assignment and analysis was performed with CARA.

Electrophoretic mobility shifts assay. Binding reactions were carried out for 30 min at room temperature in 10 μl of binding buffer (20 mM Tris pH 7.2, 80 mM NaCl, 2 mM DTT). A fixed concentration of 5'-end Cy5-labeled duplex DNA (7.5 nM) was incubated with increasing amounts of Smad4 MH1 domain. Electrophoresis was performed in non-denaturing 12.0% (19:1) polyacrylamide gels. The gels run for 1 h in 1x TG buffer (25 mM Tris, pH 8.4, 192 mM Glycine) at 90 V at 20 °C. The gels were exposed to a Typhoon imager (GE Healthcare) using a wavelength of 678/694 nm (excitation/emission maximum) for the Cy5 fluorophore.

NMR relaxation measurements. 1H, 13C, 1H-1H, 13C-13C, 1H-15N, 1H-1H-1H-1H-1H-1H-1H, 1H-1H-1H-1H-1H-1H-15N, and average fragment size of 250 bp. Additional information is described in the BMRB (entry 26945). For the titration experiments, NUS samples (500 μM for backbone and 15N-NOEY experiments and 100 μM for titrations) were equilibrated in a buffer containing 20 mM sodium phosphate and 100 mM NaCl. All samples were supplemented with 10% D2O and pH adjusted to 6.3. Spectra were acquired using 15N-labeled protein samples at the indicated concentrations with progressively increasing amounts of the unlabeled protein. Chemical shifts perturbation experiments were performed by monitoring of 15N with respect to 2H. NMRPipe and MnmDNR were used for spectroscopy and spectra assignment and analysis was performed with CARA.

Generation of CRISPR/Cas9. sgRNAs targeting genomic regions of interest were designed using CRISPR Design Tool (http://crispr.mit.edu/) and synthesized by IDT, Inc. Single cells were sorted into irradiated MEF feeder for increased viability through FACS 72 h post transfection. Mutant clones were first screened through aberrant melting temperature of qPCR products, then verified by PCR, TA cloning, Sanger sequencing, and CRISPRe-Plex analysis individually. The mESCs used for this experiment are diploid.

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two-parameter function (Eq. 1), where $I_0$ and $I(t)$ are the peak intensities at times 0 and $t$, respectively. The optimum value of the $I_0$ and the $T_1,2$ parameters are determined using the Levenberg-Marquardt optimization algorithm:

$$I(t) = I_0 e^{-t/T_1} e^{-t/T_2}$$

(1)

The rotational correlation time of the domains was calculated with the Eq. 2, using the approximation of slow molecular motion $\tau_r$ larger than 0.5 ns and assuming that only $I(0)$ and $I(\infty)$ spectral density terms contribute to the overall value. $\nuN$ is the $\nuN$ resonance frequency:

$$\tau_r = \frac{1}{\nuN} \left( \frac{T_1}{T_2} - 2 \right)$$

(2)

Heteronuclear NOE experiments were acquired in an interleaved manner (the reference and the presaturated HSQC spectra). Steady-state values of $H^1$-$^{15}N$ NOEs resulted from the ratios of the peak intensities measured in the reference ($I_0$) and the presaturated ($I_e$) spectra during the relaxation delay, as described. Background noise levels $\sigma_e$ and $\sigma_0$ were measured and used to determine the NOE standard deviation:

$$\text{NOE} = \left( \left( \frac{\sigma_e}{\tau_e} \right)^2 + \left( \frac{\sigma_0}{\tau_0} \right)^2 \right)^{1/2}$$

(3)

**SAXS data collection, analysis, and modeling.** SAXS data were collected on samples of human Smad4 MHI at protein concentrations ranging from 0.5 to 4.2 mg mL$^{-1}$, in 20 mM Tris buffer, pH 7.2, 150 mM NaCl. Data were acquired at Beamline 29 (BM29) at the European Synchrotron Radiation Facility (ESRF; Grenoble, France). Protein samples were centrifuged for 10 min at 10,000 g prior to data acquisition. Experiments on BM29 were collected at 12.5 keV and data were recorded on a Pilatus 1 M detector, at 10°C. We collected 10 frames for each of the samples (1 s exposure).

Image conversion to the 1D profile, data reduction, scaling, and buffer subtraction were done by the software pipeline available at the BM29 beamline. Further processing was done by the ATSAS software suite. Guinier plot calculation, for radius of gyration estimation, was performed by PRIMUS, included in the ATSAS suite, using low q regions ($\nuN\times\tau_r$ < 1.3).

Modeling with SAXS and NMR data were done with XPLOR-NIH, using the SAXS 1D profile and the backbone chemical shifts of Smad4 MHI. The chemical shifts were converted to backbone, phi, and psi, dihedral angles using TALOSN.

Starting from the crystal structure (PDB: 3QSV), DNA and water molecules were removed and secondary structure elements were restrained with an RMSD metric, for the determination and COOT for the manual improvement of the models. Water molecules bound at the DNA–protein interface were selected when they participate in at least three hydrogen bonds (cutoff distance of 3.5 Å).

**Ramachandran statistics.** The human Smad4–GGCGG complex has 99.2 and 0.8% of the residues in the allowed regions of the Ramachandran plot, 96.7 and 3.3% for the Trichoplax Smad4–GGCGG, 96.7 and 3.3% for the GGCT complex, 98 and 2% for the human Smad3–GGCGG complex, 98.4 and 1.6% for the Smad3–GGCT complex and 96.7 and 3.3% for Smad4–GGCGG complex. No structure displays outliers residues. The data were validated using MolProbity.

**Isotothermal titration calorimetry.** ITC measurements were performed using a nano ITC calorimeter (TA Instruments) at 20°C. DNA and protein samples were dissolved in the same buffer and degassed before the experiments. Concentrations were determined using a NanoDrop system and their predicted extinction coefficients. The NanoAnalyze software (TA Instruments) was used to analyze the binding isotherms. Baseline controls were acquired with buffer and pure DNA solutions. Fitting of the stoichiometry of the interactions, palindromic DNA nanocrystalline structure displays two theoretically equivalent sites (detected in the EMSA assays). However, if the stoichiometry is considered as one of the parameters that can be adjusted together with the $K_D$, the stoichiometry values obtained were slightly below the expected 0.5, even for the SBE used as control. We assume that these discrepancies are due to unavoidable errors in the determination of the active concentration of duplex DNA present during the titration, and also due to the equilibrium between duplex and single-stranded DNA (some duplexes have melting points close to 37 °C). Fittings were performed using the independent binding sites model. For the GCCT, we obtained $\Delta H$ of ~60 KJ mol$^{-1}$, $\Delta S$ of ~60, and $\Delta G$ of ~42. Similar values were obtained for the GGCT, ~62, ~74, and ~40, respectively. We obtained $\Delta H$ of ~24, $\Delta S$ of ~70, and $\Delta G$ of ~44 for the GGCGG.

**Computational methods.** We downloaded the mm9 and hg19 human reference genomes using the UCSC genome Distributed Annotation System server (DAS). From this, we used the EMBOSS tRNAkanitalize tool to scan for the following DNA patterns: CAGAC, AGAC, GGCCG, GGCTG, GGCGG, and GGCGC. For each of the patterns and from the output of the search, we generated a bed track that were displayed with the IGV 2.3.66 tool, for visual inspection of the results.

The following Gene Expression Omnibus (GEO) data sets were downloaded for the analysis: (GEO accession numbers ordered as shown in the figures: GSM727561, GSM727586, GSM727564, GSM727589, GSM126681, GSM727557, GSM727585, GSM539548, GSM76175722. With the exception of GSM126681, all remaining data sets were originally aligned to reference genome hg18. We have referenced them to hg19 using the UCSC liftOver tool.

We used the GEO data sets GSM727586, GSM727589, GSM727557, and GSM727585 for the human and Trichoplax Smad4 complexes. However, the results still show a notable enrichment of 5GC SBE clusters (18 and 40% of the Smad binding regions have three or more 5GC motifs vs. 6% in the baseline) in the H3K4me3 ChIP-Seq data. For these controls, we used Smad4 ChIP-Seq peaks that were not overlapped by SP1 ChIP-Seq peaks or by CTCF ChIP-Seq peaks (two different experiments) in human HepG2 cells. Each of the filtered data sets showed similar clustering values than the unfiltered Smad4 HepG2 data set. In addition, to determine whether the GC-rich regions that bind GC-rich sites affect the interaction of Smad proteins with 5GC SBEs.

**Human Smad4 GGCG complex:** the crystallization condition was 0.2 M lithium acetate, 20% PEG350. Crystals were cryoprotected in mother liquor supplemented with 20% 550 MME.

**Human Smad3 GGCGC complex:** the crystallization condition was 0.2 M lithium acetate, 20% PEG350. Crystals were cryoprotected in mother liquor supplemented with 20% 550 MME.

**Human Smad3 GGCGG complex:** the crystallization condition was 0.2 M sodium phosphate, 0.1 M BisTris propane pH 6.5, 20% PEG350 cryoprotected with 20% 550 MME.

Crystals were frozen in liquid nitrogen. Diffraction data were recorded at the ESRF on the beamlines ID23-2 (for Hu_Smad4-GGCGC and Hu_Smad4-GGGCGC), ID23-1 (for T_Smad4-GGCGC), or ID30A-3 (for Hu_Smad4-GGCT and ID29 (for Hu_Smad4-GGCGC and GGGCGG complexes). The data were processed with XDS and scaled and merged with SCALA. Initial phases were obtained by molecular replacement using PHASER from the CCP4 suite (search model PDB code: 3QSV). REFMAC5, PHENIX, and BUSTER were employed for the refinement and COOT for the manual improvement of the models. Water molecules bound at the DNA–protein interface were selected when they participate in at least three hydrogen bonds (cutoff distance of 3.5 Å).
Motif analysis. To determine each motif frequency and the corresponding average frequencies and distributions, we created 200 bp DNA regions centered in ChIP-Seq regions that were up to 1000 bp from the transcription starting sites obtaining 1755 distinct regions from Smad1 (GSM1505745), 49 from Smad3 (GSM727585), and 155 from Smad4 (GSM727586). Similar analyses were performed with the Smad4 HepG2 (ENCFF484WVM) cells data set, obtaining 16,463 peaks. We also generated a baseline data set with 500 regions of 200 bp length generated from random human genomic coordinates. We determined the number of CAGAC, AGAC, and GGCGC, GGCTG, GGCCG or GGCGG sites in both ChIP-Seq data and baseline regions.

As additional data sets, we also analyzed 200 bp DNA regions centered in ChIP-Seq peaks genome wide (not restricted to promoters) for Smad4 data sets GSM727586 (4531 regions) and ENCFEF484WVM (45,887 regions). As additional controls, we analyzed peaks in both promoters and genome wide for the Smad4 data set ENCFEF484WVM in which we discarded all peaks that had any overlap with the SPI data set ENCFEF175VXL, obtaining 6393 and 23,084 distinct regions, respectively. The same control was performed discarding overlapping peaks with the CTCF data set ENCFEF237OKO, obtaining 9035 and 36,887 distinct regions, respectively.

Data availability. Smad4 ChIP-Seq data were deposited in the Gene Expression Omnibus database under accession number GSM246361. NMR assignments and chemical shifts have been deposited in the Biological Magnetic Resonance Data Bank, BMRB entry 26945. Densities and coordinates have been deposited in the Protein Data Bank PDB entries are 5MEY, 5MEZ, SMF0, SNM9, SOAPD, and SODG76.

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Author contributions

P.M.-M. performed NMR measurements and computational analysis. M.J.M., P.M.-M., and M.B. assigned and analyzed the NMR data. Z.K. screened crystallization conditions and collected X-ray data. Z.K., R.F., M.J.M., and J.A.M. solved the X-ray structures. M.B. and M.J.M. designed the DNAs for the screenings. M.B., L.R., and A.V. performed the EMSA and ITC experiments. E.A. and L.R, cloned, expressed, and purified all proteins. T.G. collected and analyzed SAXS data. Q.W. and Y.Z. performed the CRISPR/Cas9 and Q. X. the Smad4 ChIP-Seq experiments. J.M. and M.J.M. designed the work. All authors contributed ideas to the project. J.A.M., M.J.M., and M.J.M. supervised the project and J.M. and M.J.M. wrote the manuscript.

Additional information

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