Comparative genomics reveals key gain-of-function events in Foxp3 during regulatory T cell evolution

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INTRODUCTION

Most multi-cellular organisms evolved cell-based immune mechanisms protecting themselves from invading pathogens. In higher vertebrates, the adaptive immune system maximizes its ability to recognize pathogen-derived antigens by generating a diverse repertoire of B cell and T cell receptors through the random rearrangement of gene segments (Pancer and Cooper, 2006). This recombinatorial diversification intrinsically harbors the risk of generating self-reactive specificities and thus the development of autoimmunity. The deletion of self-reactive T cells during their development in the thymus (central tolerance) has for a long time been considered the key mechanism of self-tolerance (Mathis and Benoist, 2007). However, there are non-self antigens, such as food and commensal bacteria that the immune system must tolerate (Macdonald and Monteleone, 2005). Placental pregnancy represents an even more dramatic challenge, since the maternal immune system is continually exposed to paternal antigens expressed by the fetus.

In order to tolerate the immunogenic challenges of non-self antigens, new mechanisms had to evolve and regulatory T cells (T\textsubscript{R} cells) are critical components in this type of peripheral tolerance (Fontenot et al., 2003, 2005; Horii et al., 2003; Khattri et al., 2003). These cells suppress a wide variety of undesirable immune responses (Sakaguchi, 2004) and play a crucial role in maternal–fetal tolerance (Aluvihare et al., 2004; Rowe et al., 2011).

Abbreviations: CC, coiled-coil; FKH, forkhead domain; IPEX, immunodysregulation polyendocrinopathy enteropathy X-linked syndrome; ProR, proline-rich region; T\textsubscript{H} cells, helper T cells; T\textsubscript{R} cells, regulatory T cells; ZnF, zinc finger.

The master-regulator of T\textsubscript{R} cell development and function is the lineage-specific transcription factor Foxp3 (Fontenot et al., 2003, 2005; Horii et al., 2003; Khattri et al., 2003). This protein belongs to a large family of forkhead box winged helix (Fox) transcription factors (Hannenhalli and Kaestner, 2009). Together with Foxp1, Foxp2, and Foxp4 it forms the Foxp subfamily, which is defined based on similarities in their domain architecture (Hannenhalli and Kaestner, 2009). These proteins contain several domains that are known to be of functional importance (Ziegler, 2006). The forkhead domain (FKH) is responsible for nuclear localization and DNA binding and in Foxp3 it is also responsible for interaction with the transcription factor NFAT (Lopes et al., 2006; Wu et al., 2006; Bandukwala et al., 2011). The coiled-coil (CC) is required for homotypic and heterotypic dimerization (Lopes et al., 2006) while the function of the C\textsubscript{2}H\textsubscript{2}-type zinc finger (ZnF) remains unclear.

The preponderance of evidence suggests that Foxp3 acts on multiple levels and is critical for multiple regulatory processes (Josefowicz and Rudensky, 2009). In some cases, it mediates a switch from gene activation to repression while in others it leads to enhanced transcription. Whether acting as an activator or as a repressor, it appears that Foxp3 hijacks and redirects already established transcriptional networks.

To better understand the role of Foxp3 in controlling T\textsubscript{R} cells and immune tolerance, we studied its function in an evolutionary context. We show that Foxp3 was acquired during early vertebrate evolution, but lacked key components important for its function. While the gene was retained in the lineage leading to mammals, it appears to have been lost from the genomes of birds. Our analyses...
suggest that Foxp3 evolved in a modular fashion and indicate a stepwise gain of functional domains. We found that the “first” Foxp3 ortholog that contained all of the domains required to confer T<sub>R</sub> cell phenotype can also be found in the genome of egg-laying mammals, with a further stretch of conservation gained in placentals. Our findings suggest that during its evolutionary diversification in the mammalian lineage, Foxp3 acquired several domains that allowed it to interact with a multitude of partners required for its function, making it a master-regulator of T<sub>R</sub> cell development and function.

MATERIALS AND METHODS

PROTEIN AND DNA SEQUENCES

All sequences were retrieved from the NCBI, UCSC, and Ensembl databases before January 2012. For BLAST analyses, mouse Foxp proteins were used for NCBI or Ensembl BLASTP and NCBI TBLASTN (databases: nt/nt and est_others). Specifically for Foxp3 proteins the following accession numbers were used: mouse (NP_0473380), lizard (ENSACAF00000013243; partial sequence), opossum (XP_001372400), platypus (XP_001507281/ ENSOANP00000021424), frogs (BAG12188.1 and NP_001121199.1), zebrafish (ACQ44666), salmon (NP_001185776), rainbow trout (CAT65091), stickleback (ENSGACP00000016881), and tetraodon (ENSTNIP00000007498).

SYNTENIC ANALYSIS

The UCSC genome browser was used for comparing chromosome X from mouse (9 mm; Chr X: 7,100,000–7,500,000) to chromosomes from several other species using BLASTZ alignment (Miller et al., 2007). For the construction of dotplots of genomic alignments, PipMaker (Schwartz et al., 2000) was used with default settings. For zebrafish analyses (danRer6), two loci were found containing putative Foxp3 orthologs at Chr 8: 21.9 mb and Chr 8: 24.1 mb, with the latter being the currently annotated entry in the NCBI database (GeneID: 571165). For sequence comparison in birds, the currently available chicken (galGal3) and zebra finch (taeGut1) assemblies were used. As few annotated genes can be found in the proximity of mouse Foxp3 on the centromeric side, a larger stretch of the telomeric side was used in all cases.

PROTEIN ALIGNMENT AND TREE BUILDING

Protein sequences were aligned using Muscle (Edgar, 2004) and visualized in Geneious (Drummond et al., 2010). For alignments used in phylogenetic analyses, full-length protein sequences were manually curated to remove regions with poor alignment properties. Bootstrapped trees (1000 repetitions) were constructed using Neighbor joining (substitution models: Jukes–Cantor). Dotplot protein alignments were constructed using EMBOSS dotmatcher (Rice et al., 2000) with a window size of 10 and the Blossum50 matrix. A threshold of 40 was chosen for the final output. For the conservation plot in Figure 5A, a placental consensus sequence (at least 75% identity) was created by aligning Foxp3 from mouse, rat, human, rhesus macaque, crab-eating macaque, cow, dog, and cat. A mammalian alignment was created by aligning opossum and platypus to the placental consensus and a vertebrate alignment was created by aligning frog and zebrafish. Finally, a Foxp subfamily alignment was created by aligning mouse Foxp1, Foxp2, and Foxp4 to the placental consensus. For all alignments, mouse Foxp3 was used as a reference and all alignments were trimmed so this sequence was without any gaps. For the conservation plot in Figure 5B the same mammalian alignment was used and for the non-mammalian vertebrates an alignment was created from X. tropicalis, X. laevis, rainbow trout, stickleback, tetraodon, salmon, and zebrafish. The latter alignment was manually curated to remove regions with large indels. EMBOSS plotcon was then used to plot a similarity score across the two alignments using a window size of 20.

HOMOLOGY MODELING

SWISS-MODEL (Schwede et al., 2003) was used to build homology models of Foxp3 mutated sequences, using the complex of Foxp3FKH:DNA:NFAT as a template (PDB ID: 3QRF; Bandukwala et al., 2011). The resulting structures were visualized using PyMOL (DeLano, 2002). For IPEX and signature residues, the accessible surface area was calculated with GetArea using default settings (Frazczewicz and Braun, 1998).

PREDICTION OF FUNCTIONAL DOMAINS

The coiled-coil, zinc finger and forkhead were predicted using the SMART and COILS2 programs (Lupas et al., 1991; Schultz et al., 1998). The N-terminal proline-rich (∼15% proline content) and glutamine-rich regions were determined manually. The nuclear localization signal within the FKH was determined based on the presence or absence of a RKKR motif in the FKH domain (Lopes et al., 2006).

DOMAIN SELECTION ANALYSIS

The random effects likelihood (REL) test implemented on the Datamonkey website (Pond and Frost, 2005) was used with codon-aligned Foxp3 sequences from mouse, cat, cow, macaque, dog, horse, human, pig, rat, rhesus macaque, and sheep. Prior to running the REL test, a model selection tool was executed to find the best nucleotide substitution model. All codons showing evidence of purifying selection with the REL test also replicated when we performed an independent SLAC and FEL test (Pond and Frost, 2005). All three tests agreed when using a Bayes factor cutoff value of >40.

ANTIBODIES AND FLOW CYTOMETRIC ANALYSIS

The following antibodies were used in the study (all from BD Biosciences, unless otherwise mentioned): CD3ε, CD4, CD8a, CD11b, CD11c, CD19, CD25, Ly6G, CD45RB, ratCD8a, CTLA-4, LAG-3, CD62L, I-A/I-E, and IL-2. All antibodies were used according to the manufacturer’s instructions and subsequent flow cytometric analysis was performed using a FACSCalibur system (BD Biosciences).

CONSTRUCTION OF RETROVIRAL PLASMIDS

A bicistronic MLV-based retroviral plasmid m6p8 (Bloor et al., 2008) co-expressing a GPI-linked extra-cellular part of ratCD8a (rCD8a) or m6pg co-expressing GFP, was used for all experiments. The domain-deletion mutants were created by the deletion of the following amino acids from mouse Foxp3: ΔProR:
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FIGURE 1 | Foxp3 orthologs can be found in mammals and jawed vertebrates. (A) Mouse Chr. X 710–750 mb containing the foxp3 locus and flanking regions with exons shown as vertical bars. A discontinuous whole genome nucleotide alignment is shown below as implemented in the UCSC genome browser (Kent et al., 2002). (B) Continuous dotblot genome alignments of 160 kb flanking the murine foxp3 locus with foxp3 loci of other species (black dots: aligned region; dark blue/green: forward/reverse exons; light blue/green: forward/reverse UTRs). (C) Mouse Chr. X containing foxp3 and adjacent genes with exons shown as vertical black bars. The mammalian consensus (Mammal Cons) represents 20 placental mammals available in the UCSC Genome browser with the height of the green bars denoting the level of conservation in all species determined by BLASTZ (Miller et al., 2007). A discontinuous whole genome nucleotide alignment of the mouse genes in this region with those of other species, irrespective of their location in the genome, are shown as blue bars. The height of the bars corresponds to the level of conservation. Low quality or missing sequence data is shown as gray boxes and alignment gaps are shown as gray double lines. For clarity, only regions containing genes are shown. (D) Conservation of Fox family members. The emergence of the Foxp subfamily coincides with the evolution of adaptive immunity in jawed vertebrates.

ANIMALS AND CELL PREPARATIONS
All animal care was performed by expert animal technicians in compliance with the relevant laws and institutional guidelines. All animals were age- and gender-matched Balb/c or C57/Bl6 mice, 2–4 months old and maintained under specific pathogen-free conditions. CD4+CD25+ T cells were isolated by negatively depleting lymphocytes stained with a combination of FITC-conjugated antibodies specific to CD8, CD19, CD11b, CD11c, Ly6G, and CD25.

aa1–150; ΔZnF: aa197–221; ΔCC: aa232–261; ΔFKH: aa335–429. The various Foxp3 mutations and deletions were amplified from RNA preparations using SuperScript II (Invitrogen) and KOD polymerase (Novagen) and cloned into m6p8 and m6pg. An irrelevant gene, blasticidin-S-deaminase was used as a control in all experiments. The resulting constructs were sequenced and expression of the transgenes were confirmed by flow cytometry or western-blot.

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Table 1 | Foxp Subfamily BLAST Hits.

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The Foxp1, Foxp2, Foxp3, and Foxp4 proteins from Mus musculus were used to perform standard BLAST (NCBI or Ensembl) searches and the various hits were retrieved. The table shows the accession numbers of the different hits, their length in bp, and their percent identity to mouse Foxp proteins.

**Using an AutoMACS (Miltenyi Biotec) as previously described (Andersen et al., 2008). For the preparation of antigen-presenting cells (APCs) used in suppression assays, staining with α1-A1-I-E-specific antibody was followed by positive selection using magnetic beads on the autoMACS. Subsequently the cells were irradiated (3000 rads), washed, and used directly.**

**RETROVIRAL TRANSDUCTION**

Viral supernatant was produced by transient co-transfection of 293ET cells with an equal amount of the packaging plasmid pCI-Eco and the respective m6p or m6pg constructs using standard calcium-phosphate protocols. After 48 h the supernatant was collected, filtered, and used immediately. Purified CD4+CD25− T cells were activated using plate-bound CD3ε-specific antibody (0.6 µg/ml) and recombinant mIL-2 (PeproTech; 10 U/ml). After 36 h, the cells were transduced using a 1:3 dilution of supernatant supplemented with protamine-sulfate (Sigma; 6 µg/ml), followed by centrifugation at 600 × g for 2 h at 32°C. The transduced cells were left in complete medium (RPMI/10%FCS/10 µM β-mercaptoethanol/50 µg/ml gentamicin) supplemented with recombinant mIL-2 for 48–72 h before analysis. For IL-2 intracellular staining, the cells were rested an additional 48 h and then reactivated with PMA and ionomycin in the presence of Golgi-stop (BD Bioscience) for 6 h, where after they were fixed and permeabilized using the Cytofix/Cytoperm (BD Bioscience) according to the manufacturer’s instructions.

**SUPPRESSION AND ANERGY ASSAYS**

Transduced T cells were sorted based on their co-expression of rCD8a using an AutoMACS, washed, and counted using a Vi-cell XR (Beckman Coulter) followed by flow cytometric analysis using CaliBRITE beads (BD Bioscience) as an internal standard. The purity of the sorted cells was always >95%. Target CD4+CD25− T cells were labeled with 5 µM CFSE (Molecular Probes) for 15 min, washed and counted. APCs were prepared as described above. Suppression or anergy assays were set up in U-bottom 96 well plates coated with CD3ε-specific antibody (0.6 µg/ml). About 5 × 10⁴ transduced cells were incubated together with 5 × 10⁴ CFSE-labeled target T cells and 2.5 × 10⁴ irradiated APCs. After 72 h of co-culture the cells were subjected to flow cytometric analysis. The relative proliferation of transduced T cells (anergy) and target T cells (suppression) were measured by calculating the ratio of cells/beads. For other experiments suppression by the transduced T cells was assessed by the percentage of divided target T cells calculated based on CFSE dilution.

**IN VIVO WEIGHT-LOSS ASSAYS**

Mouse CD4+CD25−CD45RBl+/− T cells were isolated from spleens of female Balb/c. Cells were enriched for CD4+ cells...
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FIGURE 2 | Syntenic analysis of the Foxp3 loci in zebrafish. (A) Mouse Chr. X 7.135–7.235 mb containing the foxp3 locus and flanking regions with exons shown as vertical black bars. A discontinuous whole genome nucleotide alignment of the zebrafish genome as determined using BLASTZ (Schwartz et al., 2003) in the UCSC genome browser (Kent et al., 2002) is shown below. Alignments from Chr. 8 are shown in red (Chr. 8 in the region around 24.1 mb), blue (Chr. 8 in the region around 21.8 mb), and green (multiple regions on Chr. 8), and those from other chromosomes in gray. (B) Continuous genome alignments of 160 kb flanking the murine foxp3 locus with the putative foxp3 locus from zebrafish on Chr. 8 in the region of 24.1 mb (black dots: aligned region; dark blue/green: forward/reverse exons; light blue/green: forward/reverse UTRs).

by MACS depletion followed by staining with PE conjugated anti-CD45RB antibody and the CD45RBhi (25% brightest) cells were purified (>99%) by FACS (MoFlo). Seven week old weight-matched female CB.17.SCID mice received an intraperitoneal (i.p.) injection of $3 \times 10^5$ CD4+CD25− CD45RBhi cells along with $2 \times 10^5$ CD4+CD25− cells transduced with either a control gene, wildtype Foxp3 or a mutant version of Foxp3 and the mice were daily monitored for weight loss.

STATISTICAL ANALYSIS

All the experiments were repeated at least twice with representative results shown in some cases. Statistical analyses were performed using Prism. Student’s t tests and one-way ANOVAs...
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FIGURE 3 | Phylogenetic relationship of Foxp subfamily orthologs. The NCBI and Ensembl databases were queried for full-length protein sequences for Foxp subfamily members. The proteins were aligned and a bootstrapped tree (1000 repetitions) was constructed with the Neighbor Joining algorithm implemented in Geneious (Drummond et al., 2010). Foxn1 was used for out-group rooting. The numbers represent bootstrap values and the scale-bar is equivalent to an average of 20% change in amino acid divergence.

were performed with \( p \) values <0.05 considered statistically significant.

RESULTS

GAIN OF FOXP3 IN JAWED VERTEBRATES

To investigate the evolutionary conservation of Foxp3 as compared to other Foxp subfamily members, we performed BLAST searches (Table 1) and detailed analyses of syntenic regions around the putative foxp3 loci (Figures 1A–C). We found that the presence of Foxp subfamily members is restricted to jawed vertebrates and foxp3-like genes (Foxp3L; 30% protein identity to mouse Foxp3) can be found in bony fish, amphibians, and reptiles, but not in birds (chicken and zebra finch). However, they are retained in the mammalian lineage, where foxp3 (>50% protein identity) is
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FIGURE 4 | Alignment of Foxp1, Foxp2, Foxp3, and Foxp4 orthologs. (A) Mouse Foxp1 amino acid (aa) sequence aligned to its orthologs in human (92.9% identity; 677aa), opossum (88.3% identity; 677aa), platypus (88.3% identity; 698aa), frog (94.8% identity; 659aa), and zebrafish (78.4% identity; 659aa). (B) Mouse Foxp2 amino acid (aa) sequence aligned to its orthologs in human (99.4% identity; 715aa), opossum (98.7% identity; 713aa), platypus (91.0% identity; 723aa), frog (94.8% identity; 738aa), and zebrafish (78.4% identity; 697aa). (C) Mouse Foxp3 protein sequence aligned to its putative orthologs in human (86.5% identity; 431aa), opossum (48% identity; 473aa), platypus (55.5% identity; 502aa), frog (34.5% identity; 693aa), and zebrafish (30.1% identity; 376aa). (D) Mouse Foxp4 protein sequence aligned to its putative orthologs in human (92.3% identity; 680aa), opossum (95.7% identity; 677aa), platypus (69.7% identity; 670aa), frog (70.8% identity; 641aa), and zebrafish (69.7% identity; 670aa). The gray arrows show the position of the putative functional domains within the aligned sequence. Identical amino acids are represented by black bars, similar amino acids by gray bars, and not similar amino acids by white bars. Gaps are shown as black line.
present in all members including marsupials and monotremes (egg-laying mammals). We found that the gene order surrounding foxp3 was highly conserved in all mammals, strongly suggesting an orthologous relationship (Figures 1A–C). The gene order surrounding the two foxp3-like genes in zebrafish was not conserved (Figures 1B and 2), but syntenic regions could be identified on
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chromosome 8 (Figures 1A and 2A). While most genes found in the vicinity of foxp3 in mammals could also be identified in birds, no foxp-like genes could be found in their proximity (Figure 1C).

In addition, phylogenetic analyses showed Foxp3 to be the most diverse member of the Foxp subfamily, especially in non-placental vertebrates (Figure 3).

Taken together, our data suggest that foxp3 or foxp3-like genes can be found in jawed vertebrates, which correlates with the presence of an adaptive immune system and thus a need for peripheral tolerance mechanisms (Figure 1D).

STEPWISE GAIN OF FUNCTIONAL DOMAINS

To gain insight into the functional potential of the identified foxp3 orthologs, we performed comparative analyses of the various proteins. We found that the sequence homology of foxp3 orthologs across all vertebrates is restricted to the C-terminal half of the protein containing the FKH, CC, and ZnF (Figure 4).

While in the other Foxp subfamily members the conservation is evenly distributed over the length of the protein (Figure 4), mammalian Foxp3 distinguishes itself from its non-mammalian counterparts by a proline-rich region (ProR; >15% prolines) spanning exons 2 and 3 (Figures 5A–C). In placentals, the homology of this region extends further into exon 1 (Figures 5A,C).

We performed a comparison of multiple sequence alignments of Foxp3 from 10 mammals with that of 5 non-mammalian vertebrates. We found that while the sequence-space occupied by the ProR in mammals is also present in non-mammalian vertebrates, the part of the N-terminal region having the best alignment to mammalian Foxp3 was chosen from these organisms.

FIGURE 6 | The ProR is disordered in placentals. (A) The disorder of Foxp3 from various species was calculated using GlobPlot (Linding, 2003). (B) The percentage of disorder in the ProR from the various species. Since not all Foxp3s have a ProR, the part of the N-terminal region having the best alignment to mouse Foxp3 was chosen from these organisms.

To test our computational predictions, we ectopically expressed wildtype Foxp3 and domain-deletion mutants lacking putative functional elements, in “normal” T cells. The forced expression of wildtype Foxp3 in non-regulatory T cells has been shown to confer a Tr cell phenotype to the transduced cells in most measurable aspects (Fontenot et al., 2003; Hori et al., 2003). The transduced cells suppress the proliferation of “target” Th cells and become anergic (non-proliferative) ex vivo. When tested in vivo, they are capable of preventing weight loss induced by co-transfer of “aggressor” cells into lymphopenic mice (Hori et al., 2003; Mottet et al., 2003). When we analyzed our domain-deletion mutants, we found that the ProR, ZnF, CC, and FKH domains all were required to confer Tr cell phenotype in vivo (Figure 7B). Ex vivo, in contrast, only cells transduced with mutants lacking the ProR or FKH appeared
FIGURE 7 | Functionally important Foxp3 domains are under purifying selection in placentals. (A) The REL (Kosakovsky Pond and Frost, 2005) test as implemented in Datamonkey (Pond and Frost, 2005) was used to perform a selection analysis of placental Foxp3s. Bayes factor scores of individual codons from the ProR, ZnF, CC, and FKH are shown (dotted red line: arbitrary Bayes factor cutoff for purifying selection). (B–E) T cells were transduced with wildtype or mutant Foxp3 as well as an irrelevant control gene. (B) The ability of the transduced cells to prevent weight loss by co-transferred “aggressor” T cells was measured by following the mean weight of the mice (n ≥ 4 per group) normalized to their weight on the day of onset in control mice. (wildtype vs. mutants: P ≤ 0.001 in each case; one-way ANOVA). (C–E) Suppressive activity and induction of anergy (non-proliferation) in the transduced T cells as shown by the relative proliferation normalized to that of control transduced cells or “target” T cells or transduced T cells. (E) Representative CFSE-profiles of “target” T cell proliferation. Error-bars represent the SEM.

to be defective in Foxp3 function. They were no longer anergic (Figure 7C) and failed to suppress target T cell proliferation (Figures 7D,E).

Together, the computational and functional tests highlight the importance of the functional domains of Foxp3 and agree with previous studies (Lopes et al., 2006; Ziegler, 2006).

MAMMALIAN EVOLUTION OF THE FORKHEAD DOMAIN
Evolutionary changes that alter the function of a highly conserved domain like the FKH are likely more subtle than the gain of an entire functional domain such as the ProR.

To gain further insight into the evolutionary differentiation of Foxp3, we investigated whether individual residues within the FKH distinguish the Foxp3 orthologs from those of other Foxp subfamily members. We found 12 such “signature” residues, 9 of which are conserved in all mammals (Figure 8A). In contrast, maximally 3 of these 12 signature residues could be found in the FKH of Foxp3L in fish and amphibians (Figure 8A).

To gain insight into the relevance of these findings, we used structural modeling to examine the effects of mutations in the signature residues and compared them to Foxp3 mutations that cause loss of TR cells in human immunodysregulation.
polyendocrinopathy enteropathy X-linked syndrome (IPEX) patients (Harbuz et al., 2010). Interestingly, the IPEX mutations target amino acids that are conserved across all vertebrates. We found that many of the signature residues may play important roles in protein–protein interaction and DNA binding (Figure 9). For example, the residues His365 and Glu401 are part of the Foxp3/NFAT interface (Figures 8B and 9H). Their reversion into the Foxp consensus residues Asn365 and Val401 is likely to negatively affect their contribution to NFAT interaction (Bandukwala et al., 2011).

The signature residues Gln376, His377, and Pro378, located directly upstream of the DNA recognition helix (H3), likely influence the way Foxp3 binds DNA (Figure 8C). While Gln376 points away from the DNA, the arginine found in other Foxp subfamily members makes close contact with the minor groove of the DNA (Figure 8C). In addition, helix 3 of Foxp3 has a substantially shorter N-terminus than helix 3 found in Foxp2 (Figure 8C). This is likely caused by the helix-breaking properties of Pro378, which in Foxp3 replaces an alanine or threonine found in the other Foxp subfamily members.

In contrast to the signature residues suggested role in DNA binding and protein interaction, we found that most IPEX mutations target residues important for FKH domain swapping and structural stability (Figures 8D and 9A–E;
To test the functional importance of IPEX and signature residues, we ectopically expressed Foxp3 mutants carrying point mutations in the various residues. We assessed their ability to confer suppressive and anergic potential and measured their influence on the regulation of the Foxp3 target genes IL-2, CD62L, LAG-3, CTLA-4, and CD25 (Hori et al., 2003; Andersen et al., 2008).

We found that the IPEX mutants strongly affected Foxp3 function; especially the mutant R397W that led to a complete loss of Foxp3 function (Figures 10C–E). Reversion of individual signature residues to that of the “non-Foxp3” consensus sequence (Figure 8A) only had minor effects on Foxp3 function (data not shown). When we replaced all signature residues predicted to be involved in NFAT interaction (E401, H365; Foxp3NFATmut), or to influence DNA binding (N376, H377, P378, I385; Foxp3DNAmut), we found them to have a marked effect in ex vivo assays (Figures 10F,G). When we tested the signature residue mutants in vivo, we found that Foxp3NFATmut had a significant effect on Foxp3 function (Figure 10H). Foxp3DNAmut also appeared to negatively influence T regulatory cell differentiation of the transduced cells in this assay, but did not reach statistical significance in our experiments (Figure 10H).
FIGURE 10 | Identification of Functionally Important Residues in the Foxp3 forkhead domain. (A) Structure of the domain-swapped FKH of Foxp3 (light green: monomer 1; dark green: monomer 2) in complex with NFAT (pink) and DNA (gray; Bandukwala et al., 2011). Red: signature residues; blue: IPEX mutations. (B) Surface accessibility of IPEX and signature residues of the domain-swapped FKH calculated using GetArea (Fraczkiewicz and Braun, 1998; *P-value < 0.05, unpaired t test). (C–E) T cells were transduced with wildtype Foxp3, IPEX mutants, or an irrelevant control gene. (C,D) Suppressive activity and induction of anergy in the transduced T cells as shown by the relative proliferation normalized to that of control transduced cells of (C) “target” T cells or (D) transduced T cells. (E) Representative FACS plots of the expression of Foxp3 target genes in the transduced T cells (at least three independent experiments). (F–H) Foxp3 mutants containing reverted signature residues predicted to be involved in NFAT interaction (NFATmut with H365N and E401V) or influence DNA binding (DNAmut with N376R, H377N, P378A, I385V) or a deletion of the entire FKH (ΔFKH) were created and expressed in T cells. As a control, the cells were transduced with an irrelevant control gene. (F) Bar graphs showing the mean fluorescence intensity (MFI) by FACS analysis of Foxp3 target genes in transduced cells as compared to the non-transduced cells within the same well (two independent experiments). (G) Suppressive activity and induction of anergy in the transduced T cells (two independent experiments). (H) The ability of the transduced cells to prevent weight loss caused by co-transferred “aggressor” T cells was measured by following the mean weight of the mice (n ≥ 4 per group) normalized to their weight on the day of onset in control mice. P values determined by one-way ANOVA; ns, not significant; *≤0.05, ***≤0.001. Error-bars represent the SEM.
Taken together, our computational and functional studies suggest that key stepwise events occurred during the evolution of Foxp3. While orthologs of the gene can be found in many higher vertebrates, functionally important characteristics such as the ProR and signature residues are unique features restricted to the mammalian lineage (Figure 11). This suggests that selection pressures specific to mammals affected the evolution of Foxp3 function and one might speculate that Foxp3+ TR cells in turn facilitated the evolution of invasive placental pregnancy.

**DISCUSSION**

Little is known about how Foxp3 evolved its ability to operate as a transcriptional repressor and activator, hereby acting as a master-regulator of TR cell development and function. Our study suggests that this happened via a stepwise gain of domains with key events occurring in the common mammalian and placental ancestors. We found foxp3-like genes in fish and amphibians, suggesting that the gene was acquired early in the vertebrate lineage. It appears that foxp3 has been lost from the genomes of birds, suggesting that at this point in evolution it may have served a non-essential or redundant role. However, the sequencing projects of birds are still incomplete and thus we cannot completely exclude the presence of foxp3-like genes.

Our analyses revealed that the C-terminal half of Foxp3 containing the CC, ZnF, and FKH is largely conserved across all vertebrates. In contrast, the ProR and the signature residues in the FKH, were novel components that appear to have been gained in the common mammalian ancestor, as they can be found in all mammals including monotremes and marsupials. In addition, the very N-terminal tail of the ProR is only conserved in placentals, suggesting a further functional differentiation.

Both the FKH and the ProR in the genomes of placentals contain signatures of purifying selection, highlighting the importance of these domains in Foxp3 function. Our data suggest that the FKH benefited from point mutations that “fine-tuned” functions such as NFAT interaction and DNA binding. These amino acids have been retained as Foxp3-specific signature residues within the mammalian lineage. Structural and functional studies support this notion (Stroud et al., 2006; Wu et al., 2006; Bandukwala et al., 2011) and demonstrate an increase in the Foxp3/NFAT contact interface over that of Foxp2, which contains the Foxp consensus residues in this region (Bandukwala et al., 2011).

The acquisition of the ProR in mammals may have increased Foxp3’s evolutionary differentiation from other Foxp proteins by allowing it to bind to factors involved in T cell activation and lineage commitment. Foxp3 interacts with a vast array of proteins via this domain, including the DNA binding factor EOS (Pan et al., 2009), the histone deacetylases HDAC7/9, the histone acetyltransferase Tip60 (Li et al., 2007), as well as the transcription factors RORγt (Ichiyama et al., 2008; Zhou et al., 2008), RORα (Du et al., 2008), and c-Rel (Loizou et al., 2011). It is plausible that the ability of Foxp3 to bind to these factors might have significantly increased its capacity to control TR cell development and function in the mammalian lineage.

It has been suggested that foxp3-like genes that can be found in fish are expressed in the lymphocyte lineage and fulfill similar T cell specific functions (Quintana et al., 2010; Wen et al., 2011). However, as these divergent orthologs appear to lack key features of mammalian Foxp3, further studies need to address to what extent Foxp3L can re-program cells in non-mammalian vertebrates. While Foxp3L appears to confer some suppressive activity (Quintana et al., 2010; Wen et al., 2011), the reprogramming of T cells by Foxp3 in mammals is likely to be far more widespread and affect many characteristics of TR cell function.

In conclusion our data suggest that Foxp3 gained the ability to interact with multiple partners in a stepwise fashion. These interactions can occur individually or as a complex (Li et al., 2006). Although it is impossible to directly prove that there is a link between the evolution of TR cells and that of invasive placenta- tion, our findings are consistent with this hypothesis. Based on the conservation of Foxp3, it appears that the common mammalian
ancestor had T<sub>R</sub> cells with similar characteristics to those in mice and humans. The evolution of CD4<sup>+</sup>T cell memory, which is believed to have occurred in the mammalian lineage rendered negative selection inadequate for the suppression of undesirable immune responses (Lane et al., 2010). This mechanistic niche appears to have been filled by the evolution of T<sub>R</sub> cells which provided nature with a mechanism to suppress undesirable immune responses in a localized and specific fashion (Gaspal et al., 2011). This in turn is likely to have been required prior to the evolution of invasive placentation. Once acquired, Foxp3 was subjected to strong purifying selection in placentals, consistent with the pivotal role of T<sub>R</sub> cells in maternal–fetal tolerance (Aluvihare et al., 2004; Rowe et al., 2011).

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