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Highlights

- The lengths of two polyQ repeats (Q₁ and Q₂) in FOXP2 vary broadly across mammals.
- The Q₁/Q₂ length ratio correlates quantitatively with vocalization frequency ranges.
- The FOXP2 polyQ repeats form α-helical coiled coil structures and promote LLPS.
- Evolutionary polyQ length variation regulates FOXP2 structure, LLPS, and function.

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PolyQ length-based molecular encoding of vocalization frequency in FOXP2

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SUMMARY
The transcription factor FOXP2, a regulator of vocalization- and speech/language-related phenotypes, contains two long polyQ repeats (Q1 and Q2) displaying marked, still enigmatic length variation across mammals. We found that the Q1/Q2 length ratio quantitatively encodes vocalization frequency ranges, from the infrasonic to the ultrasonic, displaying striking convergent evolution patterns. Thus, species emitting ultrasonic vocalizations converge with bats in having a low ratio, whereas species vocalizing in the low-frequency/infrasonic range converge with elephants and whales, which have higher ratios. Similar, taxon-specific patterns were observed for the FOXP2-related protein FOXP1. At the molecular level, we observed that the FOXP2 polyQ tracts form coiled coils, assembling into condensates and fibrils, and drive liquid-liquid phase separation (LLPS). By integrating evolutionary and molecular analyses, we found that polyQ length variation related to vocalization frequency impacts FOXP2 structure, LLPS, and transcriptional activity, thus defining a novel form of polyQ length-based molecular encoding of vocalization frequency.

INTRODUCTION
Vocalization is a biological process based on the emission of sound through the respiratory system underlying various forms of animal communication, e.g., human speech and oral language, intra-/interspecific signaling, as well as echolocation-based spatial navigation and prey hunting. Vocalizations can be emitted in the sonic frequency range, as defined by the human hearing sensitivity (~20–20,000 Hz), and/or in the infrasonic (<20 Hz) or ultrasonic (>20 kHz) ranges. Vocalization and hearing frequency ranges generally co-evolve to allow efficient intraspecific communication. Thus, the ability of mammals to vocalize in frequency ranges inaccessible to other vertebrates arose also in relation with key evolutionary innovations in their hearing systems.

The genetic and molecular bases of vocalization are only partially understood. In particular, it is unclear how specific features of vocalization, such as its frequency range in each species, are encoded at the genetic and molecular levels. Some genes related to vocalization, speech, language, and their evolution, have been identified and, among these, FOXP2 has been the most intensively studied. FOXP2 and its paralog gene FOXP1, both encoding for transcription factors, are involved in neural development and function, and are expressed in areas of the nervous system and sensory organs related to the emission and auditory processing of vocalizations (e.g., cerebral cortex, basal ganglia, auditory pathways, cochlea). These two transcription factors are also involved in the development of non-neural structures relevant to vocalization, such as the lung and the craniofacial skeleton.

Mutations in both genes can give rise to various forms of speech and language impairments in humans. FOXP2 mutations can cause the ‘FOXP2-related speech and language disorder’ (FOXP2-SLD) with childhood apraxia of speech (CAS) as a core phenotype. FOXP2-SLD can also include oral-motor dyspraxia, dysarthria, moderate-to-severe receptive and expressive language disorder, reading impairments, and fine motor difficulties. Large 7q31 deletions or reciprocal-balance translocations disrupting FOXP2 together with neighboring genes give rise to more complex phenotypes (‘FOXP2-plus-related disorder’). While hearing is not...
implicated in FOXP2-SLD, the possibility exists that higher-level auditory processing may be affected, as suggested by studies in genetically modified animal models. The molecular evolution of FOXP2 has also been associated with the emergence of specialized vocalization phenotypes. Indeed, two amino acid substitutions in human FOXP2, which are not present in the great apes, and intronic variants, have been related to the acquisition of speech and language. Furthermore, accelerated molecular evolution of FOXP2 has been detected in taxa with highly differentiated vocalization/hearing phenotypes related to echolocation, such as Chiroptera. Nevertheless, the specific impact of most of the observed evolutionary FOXP2 mutations on molecular and organismal phenotypes are still largely unclear.

Interestingly, both the FOXP2 and FOXP1 proteins contain two major polyglutamine (polyQ) repeats. One of these in FOXP2 represents the longest polyQ repeat in the human proteome. PolyQ tracts can form α-helical coiled coils (CCs) that mediate protein oligo-/poly-merization and interactions, regulating the activity of transcription factors in a length-dependent manner. PolyQ CC regions, as CC domains in general, can drive the formation of functional molecular condensates through liquid-liquid phase separation (LLPS), as found for RUNX2, which bears polyQ repeats forming CCs. However, the structural and functional roles of these long repeats in FOX2P and FOXP1 are essentially unknown.

The evolutionary variation of polyQ and other amino acid repeats, in terms of occurrence and length, can have an evolutionary impact by adjusting, as ‘tuning knobs’, the transcriptional activity of developmental proteins. Thus, quantitative analyses of amino acid repeats, in terms of occurrence rate, length, and length ratios, can help characterize molecular evolution processes. The full range of evolutionary polyQ length variation in FOXP2, and its possible structural and functional impact, are not yet defined.

Vocalization and hearing frequency ranges varied extensively throughout phylogeny, and similar specialized forms of vocalization/hearing developed in different branches of the mammalian evolutionary tree by convergent evolution. Thus, high- and low-frequency specialist species, vocalizing/hearing in the ultra- or infra-sonic ranges, respectively, are found in multiple orders including Primates, Rodentia, Chiroptera, Cetartiodactyla, and Afrotheria. Convergent evolution of organismal phenotypes between distant species/taxa can arise from convergent changes at the molecular level. For instance, in chiropteran and cetacean species using ultrasonic echolocation, prestin and other hearing-related proteins display similar amino acid substitutions which regulate protein function. However, none of these convergent mutations is common to all high-frequency specialist species as convergence at the organismal level may arise at the molecular level from multiple mutational pathways in partially overlapping sets of genes/proteins.

Limited knowledge exists on convergent molecular evolution in vocalization-related proteins. The order Chiroptera comprises the Yinpterochiroptera and Yangochiroptera suborders as a model taxon characterized by species with highly diversified ultrasonic vocalizations (USVs). Primary sequence analyses showed considerable length variation of FOXP2 polyQ repeats in Chiroptera in comparison with other taxa like Primates. Thus, we reasoned that it would be easier in this taxon to reveal relations between polyQ variation and vocalization-related phenotypes. Indeed, the two major N-terminal polyQ repeats, i.e., Q1 and Q2 (40 and 10 Qs in humans, respectively), display relatively limited variation in Primates. Conversely, Q1 and Q2 vary extensively across chiropteran species, in terms of both total (Q1+Q2 sum) and relative (Q1/Q2 ratio) lengths.

Notably, several species in both taxa exhibit a recurrence of interspersed proline residues and/or polyproline stretches in Q1, which shorten this repeat compared to other related species. The order Chiroptera comprises the Yinpterochiroptera and Yangochiroptera suborders. ‘Microbats’ emitting laryngeal USVs, with constant frequency (CF) or frequency modulation (FM) and variable bandwidth/harmonics, are found in both suborders.
Figure 1. FOXP2 polyQ lengths encode ultrasonic vocalization frequency in Chiroptera

(A) Upper panel: Scheme of the primary sequence of the human FOXP2 protein. The two major polyQ repeats (Q1 and Q2) are highlighted in red within a broader Q-rich N-terminal region in pink. Known domains are in light gray, i.e., zf: Zinc-finger, lz: leucine zipper; FH: Fork-head DNA binding domain. Residues N303 and S325, distinguishing the human ortholog among Primates, are in shades of green. Lower panel: partial sequence alignments of the polyQ regions (left) and of the region flanking the 303/325 positions of the human FOXP2 ortholog (right) in representative species of Primates and Chiroptera. Species names are abbreviated using the first three letters of genus and species, as reported in Table S1. On the left, unscaled phylogenetic trees (from TimeTree.org). Note how polyQ lengths vary also by proline residues insertion in Q1 and how certain bats bear human-like or other substitutions at positions corresponding to 303/325 in the human ortholog.

(B) On the left, phylogenetic tree of the chiropteran species that were analyzed from TimeTree.org (species names in Table S1). On the right, the taxon bars specify for each species the suborder (SO; Yinpterochiroptera, YIN, and Yangochiroptera, YANG), and superfamily (SF) or family (F). The call panel indicates...
For simplicity, we group FM USVs into ‘FM1’ (single harmonic, or with peak harmonic as the fundamental; Collen\(^55\) types 3, 7–8, 10), and ‘FM2’ (multi-harmonic; types 4–6, 9). Yangochiroptera also include ‘megabats’ that do not echolocate (“NE’) using laryngeal USVs but navigate relying on sight and rudimentary echolocation, using sonic wing- or tongue-generated clicks.\(^57\)

Notably, the use of these different USV types is not strictly determined by phylogeny in Chiroptera.\(^55\) Although specific USV types, such as CF in Rhinolophoidea, FM\(_1\) in Vespertilionoidea, and FM\(_2\) in Noctilionoidea, are prevalent in each chiropteran superfamily, several species demonstrate phenotypic convergence by employing USV types distinct from those of their closest relatives (as highlighted by arrowheads in Figure 1B). For example, Megaderma spasma (Rhinolophoidea) emits FM calls, Pteronotus pereirai (Noctilionoidea) CF calls, and Mormoops blainvillii (Noctilionoidea) uses FM\(_2\) calls. USV-emitting bats exhibit morpho-functional specializations, including an increased relative cochlear size\(^55\) (Figures 1C and S1A) and height (Figure S1B).

We measured Q\(_1\) and Q\(_2\) lengths in chiropteran FOXP2 orthologs, calculated their total (Q\(_1+Q_2\) sum) and relative (Q\(_1/Q_2\) ratio) lengths, and studied their variation in relation to phylogeny and USV types (Figure 1B, right columns). This analysis corroborated the initial evidence of generalized polyQ variation in Chiroptera. A comparable analysis was conducted for two analogous polyQ repeats (‘Q\(_1\)’ and ‘Q\(_2\)’ in FOXP1, with length variation being restricted to only a few species (Figure S1C).

If the observed polyQ variation in FOXP2 is related to USV, we reasoned that polyQ length may correlate with vocalization-/hearing-related parameters. Thus, USV-emitting species were divided in groups based on USV frequency in 10 kHz bins. Then, we calculated the mean Q\(_1+Q_2\) and Q\(_1/Q_2\) for each group and correlated them with the respective mean USV frequencies. The source of USV frequencies was an extensive standardized dataset provided by Collen\(^55\) which includes data for more than 300 bat species.

Notably, both the polyQ sum and ratio exhibit a strong correlation with the USV characteristic frequency (r = −0.97 for Q\(_1/Q_2\), n = 8 groups, p < 0.0001; Figure 1D). Similar correlations are observed with minimum frequency and bandwidth (r = −0.90 and 0.81 for Q\(_1/Q_2\), respectively, n = 8−9, p < 0.01 in both cases; Figure 1D). Additionally, a hearing-related morphological parameter, the relative cochlear size (calculated as the ratio between cochlear and basicranial widths, CW/BW, using data from Simmons et al.\(^56\)) also shows significant correlations with polyQ repeat lengths. When species, including NE, are grouped based on cochlear size, the mean CW/BW ratio inversely correlates with the mean Q\(_1+Q_2\) sum (r = −0.75, n = 8, p < 0.03; Figure 1D).

In summary, these findings indicate that the observed polyQ length variations in Chiroptera are closely related to vocalization- and hearing-related parameters, thus encoding USV frequency quantitatively.

**Convergent polyQ evolution in FOXP2 in relation to vocalization type in Chiroptera**

The total and relative polyQ lengths exhibit differential variations among chiropteran species with different vocalization types (CF, FM, and NE; Figure 1B). This prompted us to investigate whether polyQ variation might be related not only to USV frequency but also to USV type. Therefore, we tested whether a combination of the two polyQ-related parameters could predict vocalization type by generating a scatterplot of polyQ sum vs. ratio (Figure 1E).

Remarkably, this graph demonstrates a clear clustering of the 53 species that were analyzed into four distinct groups based on their USV types (CF, FM\(_1\), FM\(_2\), NE) rather than their phylogenetic relationships (Figure 1F), with only two exceptions. The NE species are identified by a polyQ sum >60, while the USV-emitting species (sum <60) are further divided into three clusters based on their polyQ ratio (less than 1 for the CF group, 1–2.1 in the FM\(_2\) group, and from 2.1 to ~3.5 in the FM\(_1\) group). The threshold values for these groups are highlighted in Figure S1D.

Indeed, it is worth noting how species belonging to the same Yinpterochiroptera suborder are differentially distributed among all four clusters based on their USV types. Specifically, Pteropodidae (NE) and Rhinolophoidea species utilizing CF USVs form two distinct clusters, well separated from those of FM species. Strikingly, the three Rhinolophoidea species emitting FM rather than CF USVs (M. spasma, Megaderma lyra, and Craseonycteris thonglongyai), cluster together with Yangochiroptera species that also emit FM calls. Similar cases of
Figure 2. FOXP2 and FOXP1 polyQ lengths correlate with vocalization/hearing-related parameters in Euarchontoglires

(A) As in Figure 1B, but for the superorder Euarchontoglires (species names as in Table S1). SO in the taxon bars indicates here ‘suborder’. Silhouettes of representative species are reported on the right.

(B) As in Figure 1A, partial alignments of the polyQ repeat regions of FOXP2 (upper) and FOXP1 (lower) of the indicated species of Euarchontoglires and representative chiropteran species. Taxa and USV-emitting species are highlighted by symbols according to the legend in the inset. Note, for FOXP2, the polyQ convergence with USV-emitting bats (Rhi fer, Meg spa) of multiple Euarchontoglires species known to have ultrasonic vocalization/hearing, through
convergence can be observed across the FM 1 and FM 2 clusters. Mormoops blainvillei (Noctilioidea) clusters with FM 1 Vespertilionoidea species, while Molossus molossus (Vespertilionoidea) clusters with FM 2 Noctilionoidea. Taphozous melanopogon (Emballonuridae) groups with other FM 3 species belonging to Noctilionoidea and Rhinolophoidea.

Two exceptions to the generalized clustering based on USV types are observed. First, Pipistrellus pipistrellus, an FM 1 bat, shows convergence with FM 2 bats. This convergence is of uncertain significance as it falls within the FM range and may be related to vocal plasticity in this species. The second exception is related to the CF species Noctilio leporinus and Pteronotus parnellii (Noctilionoidea) that do not converge with CF Rhinolophoidea. However, the lengths of FOXP1 polyQ repeats in both species vary considerably compared to most chiropteran species. Noticeably, this is also observed in other species that demonstrate USV type convergence with those from different orders (Figure S1C). The calls are multiharmonic and, as shown for one of them in detail in the upper panels, they extend in the ultrasonic range, well above 20 kHz (red line).

Convergent polyQ evolution related to high-frequency vocalization/hearing in Euarchontoglires

We further tested whether our findings in Chiroptera can also be generalized to other mammalian taxa, starting with Euarchontoglires (~50% of mammalian species). The superorder comprising Primates, Rodentia, and Lagomorpha, in which numerous species are able to emit USVs (Tables S1,S2, S3, and S4). FOXP2 ortholog sequence alignments revealed a striking recurrence of the polyQ variation patterns and proline insertions, as found in USV-emitting bats, also in Euarchontoglires species with known ultrasonic vocalization/hearing (Figures 2A, S2A, and S2B). In these species, Q1/Q2 is < 3.5 (as in P. pannellii) or ≥ 2 (as in N. leporinus) due to prevalent Q2 or Q1 elongation, respectively (Figure S1C). Moreover, Pteronotus pannellii also exhibits a unique proline substitution within the highly conserved FOXP2 hinge between Q1 and Q2 (Figure S1E), further distinguishing it from Noctilionoidea (see discussion).

Overall, these findings show how FOXP2 polyQ lengths can largely predict USV type in Chiroptera. Moreover, they highlight the parallelism between phenotypic USV type convergence and molecular-level polyQ length convergence in FOXP2, with a few exceptions where enhanced polyQ variation is observed in FOXP1.
Sum-vs-ratio scatterplots (Figure 2C) highlight these recurring patterns of polyQ convergence. In the FOXP2 scatterplot, multiple species with known high-frequency vocalization/hearing fall into the region occupied by USV-emitting bats (sum <60, ratio <3.5; Figure 2C, red boundaries).

In Primates, three USV-emitting species, i.e., Carlito syrichta, Microcebus murinus, and Callithrix jaccus, display polyQ convergence with USV-emitting bats. Notably, C. syrichta, the only Primate known to emit purely ultrasonic calls,\(^6^{66}\) has the lowest polyQ ratio. In Dermoptera, Galeopterus variegatus, whose ultrasonic vocalizations have been recently demonstrated,\(^6^{67}\) also displays a Q1/Q2 ratio below 3.5.

In Rodentia, several species have FOXP2 polyQ lengths in the range of USV-emitting bats, such as Microtus ochrogaster,\(^6^{68}\) Octodon degus,\(^6^{69}\) and Cricetus griseus.\(^7^{70}\) Strikingly, these species include Typhlomys cinereus, a blind rodent recently discovered to rely on echolocation for spatial navigation\(^7^{71}\) and showing molecular convergence with bats also for the hearing-related prestin protein.\(^6^{68}\)

Nevertheless, numerous USV-emitting rodents, including the mouse and the rat, display no convergence with bats. Remarkably, however, Rodentia show overall considerable variation of the FOXP1 polyQ repeats, converging with USV-emitting bats like P. pamelii and M. lyra in having a low FOXP1 Q1/Q2 (<0.5). This ratio goes below 0.2 upon extreme Q2 expansion, as evident in some species with the highest vocalization/hearing frequencies, such as the mouse, the rat and the echolocating species T. cinereus. Species like Jaculus jaculus and Marmota marmota, adapted to desert and/or fossorial life, which are sensitive to both low-frequency and ultrasonic acoustic signals with a bimodal frequency sensitivity,\(^6^{69},7^{70}\) have an intermediate Q1/Q2 ratio (0.2–0.5; also see below Afrotheria; Figures S4C and S4D).

All the analyzed Lagomorpha species have FOXP2 repeats in the range of USV-emitting bats. Two leporids (Oryctolagus cuniculus and Lepus europaeus) also converge with some USV-emitting bats in having a FOXP1 Q1/Q2 ≥ 2 (Figure 2D). While our knowledge on Lagomorpha vocalization/hearing is fragmentary, our findings are consistent with Oryctolagus audiograms showing ultrasonic sensitivity\(^7^{71}\) and recent reports of USVs in Ochotona daurica.\(^6^{61}\)

The pika Ochotona princeps (Och pri) is an extreme case of low polyQ sum and ratio, clustering with USV-emitting bats. Strikingly, O. princeps displays accelerated evolution in the hearing-related protein prestin similar to echolocating bats,\(^7^{72}\) strongly suggesting its ability to vocalize/hear in the ultrasonic range.

To support this hypothesis as a case study of the predictive power of FOXP2 polyQ lengths for vocalization/hearing phenotypes, we analyzed vocalization recordings of O. princeps in the wild and studied its inner ear. We found evidence of spontaneous calls extending into the ultrasonic range, well above 20 kHz (>25 kHz; Figure 2D; see STAR methods). Moreover, available μ-CT scans revealed, in comparison with other Lagomorpha, an even larger, higher cochlea, within a highly pneumatized temporal bone (Figure 2E), consistent with high-frequency hearing sensitivity.\(^7^{73}\)

Overall, these findings indicate that the observed convergence patterns of FOXP2 and FOXP1 polyQ repeats in relation to vocalization/hearing frequency are not unique to Chiroptera and can be generalized to other orders within Euarchontoglires.

**PolyQ lengths correlate with hearing frequency-related parameters in Euarchontoglires**

As for Chiroptera, we also searched for quantitative relations between polyQ lengths and vocalization/hearing-related parameters in major Euarchontoglires orders. Given the previous findings, we performed this analysis in Primates and Lagomorpha for FOXP2 and in Rodentia for FOXP1.

As standardized measurements of vocalization-/hearing-related parameters for many Primates and Lagomorpha species are not available, we directly measured morphological cochlear parameters related to hearing frequency in species of interest (Figures 2F, 2G, and S2E–S2G).

Thus, we determined the relative cochlear size (CW/BW) and height (height/width)\(^6^{62}\) in representative species or genera of Primates (n = 10), Lagomorpha (n = 4) and, for comparison, Chiroptera (n = 16) (Figure S2E). This analysis revealed that cochlear parameters in these taxa correlate with polyQ lengths in FOXP2 (Figure 2F). Indeed, the Q1/Q2 ratio inversely correlates with relative cochlear size and height (r = −0.72 and −0.60, respectively, n = 26, p < 0.001 in both instances; Figures 2G and S2F). Thus, species with a lower ratio have significantly larger and higher cochleae (Figure S2G).

In Rodentia, we analyzed the correlation between FOXP1 polyQ lengths and a functional hearing-related parameter, the maximum audible frequency, in a sample of species with both parameters available. We found again an inverse correlation by which species with a low ratio, like M. musculus, tend to have auditory sensitivity to higher frequencies (r = −0.64, n = 11, p < 0.04; Figure 2H).

Taken together, these findings show that polyQ length variation in FOXP2 and FOXP1 encodes vocalization/hearing frequency-related parameters also in non-chiropteran mammalian taxa.

**Convergent polyQ evolution related to low-frequency/infrasonic vocalization in mammals**

Thus, after Chiroptera and Euarchontoglires (~70% of mammalian species),\(^5^{54}\) we completed our analyses with the remaining placental super-orders, i.e., non-chiropteran Laurasiatheria (Carnivora, Pholidota, Perissodactyla, Cetartiodactyla, Eulipotyphla), Xenarthra, and Afrotheria (Figures 3A and S3; see Figure S3A for the few non-Eutherian species that could be analyzed). These taxa are of particular interest as they contain multiple species characterized by ultrasonic or infrasonic vocalization/hearing. For instance, insectivorans in Laurasiatheria (Eulipotyphla) and Afrotheria (Afrosoricida) independently developed ultrasonic signaling,\(^6^{65}\) while species with high body mass in multiple taxa, such as whales and hippos (Cetartiodactyla), elephants (Afrotheria), and rhinos (Perissodactyla) convergently evolved infrasonic vocalization/hearing.\(^4^{47},7^{77}\)

Strikingly, these two patterns of evolutionary phenotypic convergence toward either ultrasonic or infrasonic vocalization/hearing are paralleled, at the molecular level, by FOXP2 polyQ variations of opposite sign (Figures 3B and 3C).
Figure 3. FOXP2 polyQ repeat lengths quantitatively encode vocalization frequency from the infrasonic to the ultrasonic range in Mammalia

(A) As Figure 1B, for species belonging to non-chiropteran Laurasiatheria orders (i.e., Carnivora in red, Pholidota in yellow, Perissodactyla in light green, Cetartiodactyla in dark green, Eulipotyphla in cerulean), Xenarthra (light blue), and Afrotheria (blue). 

(B) As in Figure 1A, partial alignments of the FOXP2 polyQ regions for the indicated species belonging to phylogenetic tree shown in panel A. Taxa are highlighted by symbols according to the legend in the inset. The human sequence (Q1/Q2 = 4) is shown in comparison with species with known ultrasonic (Q1/Q2 < 3.5; high-frequency sinusoid) or infrasonic (Q1/Q2 < 4.5; low-frequency sinusoid) vocalization/hearing. The first group comprises some insectivorans (Microgale talazaci, Mic tat, Afrotheria, Erinaceus europaeus, En eur, Eulipotyphla) and laurasiatherian Carnivora (Felix catus, Fel cat¹); Neovison vison, Neo vis²; Enhydra lutris, Enh lut;⁷⁶ . The second group comprises Hippopotamus amphibius, Hip amp, Cetartiodactyla; Ceratotherium simum, Cer sim, Perissodactyla; Elephas maximus, Ele max, Afrotheria, Balaenoptera musculus, Bal mus, Cetartiodactyla; Physeter catodon, Phy cat, Cetartiodactyla. See results for Xenartra (Tamandua tetradactyla, Tam tet, Dasypus novemcinctus, Das nov). The vertical red line marks the Q1+Q2 threshold of ~60 beyond which only NE bats and Heterocephalus glaber are found. 

(C) Scatterplot as in Figure 2C, for the FOXP2 orthologs of species shown in panel A. The background red and blue gradients highlight the portions of the graph occupied by species with high- or low-frequency vocalization/hearing, respectively. Three species are omitted for clarity (Manis javanica and M. pentadactyla, both with Q1+Q2 = 44, Q1/Q2 = 10, and Trichechus manatus, with Q1+Q2 = 59 and Q1/Q2 = 8.83; see Figure S3). 

(D and E) Scatterplots with regression lines (in red) highlighting significant correlations between mean FOXP2 Q1/Q2 ratio and mean minimum (D) or maximum (E) vocalization frequency in groups of mammalian species binned by the same two parameters (see STAR methods). 

(F) Scatterplots with regression line (in red) display the correlation between FOXP2 Q1/Q2 ratio and log body mass (g) in 107 species of mammals with both parameters available. 

(G) Bar graphs plotting the mean log minimum (left) and maximum (middle; without Cetacea) vocalization frequencies, and the mean log body mass (right) in groups of species with a FOXP2 Q1/Q2 ≤ 3.5 (red), between 3.5 and 4.5 (gray), and >4.5 (blue). Data are represented as mean ± SEM, and the statistical tests that were performed are reported in the results section. Asterisks indicate statistically significant differences. See also Figures S3 and S4 and Tables S1, S2, and S3.

Thus, species with known ultrasonic vocalization and/or hearing⁴⁵,⁷⁴–⁷⁶ from Carnivora (e.g., Felis catus, Neovison vison, and Enhydra lutris), Eulipotyphla (e.g., Erinaceus europaeus) and Afrotheria (Microgale talazaci) have Q1/Q2 ≤ 3.5 (Figure 3B). Interestingly, also in these species Q1 shortening can arise from proline/polyP insertions. In sum-vs-ratio scatterplots these species fall in the region of USV-emitting Chiroptera and Euarchontoglires (Q1+Q2 ≤ 60; Q1/Q2 ≤ 3.5; Figure 3C).

Conversely, large species that evolved infrasonic vocalization/hearing⁷⁷,⁷⁸,⁷⁹ such as Balaenoptera musculus (Mysticeti, Cetacea), Elephas maximus (Afrotheria), and Ceratotherium simum (Perissodactyla) all have Q1/Q2 > 4.5 (Figure 3B), occupying the upper part of the scatterplot above Homo (Q1/Q2 = 4) and other species with sonic vocalization/hearing ranges (Figure 3C). Hippopotamus amphibius (Cetartiodactyla) is a borderline case (Q1/Q2 = 4.38). Other large animals in Artiodactyla/Perissodactyla, also had Q1/Q2 > 4, except for some camelsids, consistent with their ability to hear ultrasonic frequencies⁸⁰ and emit the highest frequency vocalizations in Artiodactyla.⁸¹ 

Notably, Odontoceti cetaceans such as Physeter catodon and Tursiops truncatus, which use (ultra)sonic clicks for underwater echolocation, do not cluster with USV-emitting bats, but have FOXP2 polyQ repeats like infrasonic Mysticeti (ratio >4.5; Figure 3C; a few predicted sequences also bear a possibly spurious extension of the hinge peptide, see STAR methods). This finding parallels the lack of FOXP2 convergence between USV-emitting Chiroptera and Cetacea in terms of single residue substitutions,⁸² consistent with distinct vocalization systems. Indeed, while Chiroptera produce laryngeal vocalizations, Odontoceti emit non-laryngeal clicks through the forehead not requiring orofacial coordination, a process involving FOXP2⁸³ (see discussion). The position of Odontoceti near Mysticeti in the scatterplot seems thus related to minimum, rather than maximum (USV-related) vocalization frequencies.⁸⁴,⁸⁵ 

Vocalization and hearing abilities are scarcely characterized in most Xenarthra species (Figure 3C). Interestingly, however, the fossorial Dasypus novemcinctus, thought to have low-frequency hearing,⁸⁶,⁸⁷ occupies the upper part of the scatterplot (ratio >4.5) while the insectivoran Tamandua tetrapoda is in the lower part, together with other non-xenarthran insectivorans. 

In all these taxa, some species known to produce USVs also converge with some USV-emitting bats and most rodents in having a low FOXP1 Q1/Q2 ratio (<0.5; Figure S3C). These include only two insectivoran species from Eulipotyphla (i.e., Erinaceus europaeus and Sorex Araneus)⁸⁸,⁸⁹ and all of the species from Afrotheria. Thus, Afrotheria and Rodentia together constitute the vast majority of the species with a FOXP1 Q1/Q2 ratio <0.5. 

Overall, these findings show that FOXP2 polyQ lengths vary and converge in opposite ways in mammalian species with high versus low frequency laryngeal vocalization/hearing, in the entire range from the infra- to the ultra-sonic. Interestingly, some USV-emitting species such as M. lyra (Chiroptera) and T. cinereus (Rodentia) show convergence for both FOXP2 and FOXP1. 

The FOXP2 Q1/Q2 ratio quantitatively encodes vocalization frequency ranges in Mammalia

The previous findings suggested that polyQ parameters may overall encode vocalization frequency ranges in mammals, and we tested this hypothesis for a sample of >100 species with available data. We divided these species in ten groups based on increasing Q1/Q2 ratio (in 0.5 unit bins) and calculated for each group the mean minimum and maximum vocalization frequency as found in the literature (Figures 3D and 3E). We used in our correlation analyses the minimum/fundamental and maximum frequencies as overall indicators of the frequency range in each species, which can be very wide. These vocalization frequency parameters were derived from available datasets containing information for a large number of species (see Table S2 and STAR methods), rather than from a collection of studies in single species, to avoid data selection biases when discordant frequency values are reported by different sources. We found a strong significant correlation between mean Q1/Q2 and mean minimum (r = 0.93) or maximum
Figure 4. The FOXP2 polyQ region forms \( \alpha \)-helical CCs with length-dependent stability

(A) Primary sequence alignment of four peptides comprising the FOXP2 polyQ repeats, the hinge peptide, and a few flanking residues in the Homo sapiens (Hom sap), Cynopterus sphinx (Cyn sph), Miniopterus schreibersii (Min sch), and Rhinolophus fernenueimun (Rhi fer) orthologs. Glutamine (Q) proline (P) residues are highlighted in red and blue, respectively, other residues are in black.

(B) Secondary and supersecondary structure predictions for the Q-rich N-terminal domains (NTDs; first ~300 residues) of the FOXP2 orthologs of the species listed above for panel A. On top, the thin vertical bars mark glutamine (red) and proline (blue) residues along the sequence. The gray bars highlight the regions predicted by PSI-PRED to form \( \alpha \)-helical structures. The middle panels display the CC propensity calculated by Parcoil2\(^{36,37} \) and expressed as 1-P-score, as in Fiumara et al.\(^{37} \) Regions with high CC-propensity (0.8–1) are highlighted in black (peaks cc1–cc3). Red arrows highlight, in comparison with Hom sap, cc3 elongation related to Q2 expansion in Cyn sph, cc2 shortening related to Q1 contraction in Min sch, and further cc2 shortening by Q1 interruption caused by numerous proline residues in Rhi fer, which greatly expand the gap between cc2 and cc3. Below, details of Raptor-X atomic-level structural predictions highlighting the hairpin structure formed by Q1 and Q2 \( \alpha \)-helices separated by a turn at the level of the hinge peptide. Glutamines are in red, prolines in blue, and Q-Q hydrogen bonds between the Q1 and Q2 are in yellow. Note how longer polyQ tracts correspond to longer \( \alpha \)-helices and how the loop between the two helices is greatly extended by proline insertions in Rhi fer.

(C) Lateral (left panel) and zenithal (right panel) views of Raptor-X structural model of the Cyn sph NTD region encompassing Q1, Q2, and the flanking Q-rich regions. Note how the supercoiled \( \alpha \)-helical CC hairpin extends through the Q-rich regions.

(D) CD spectra for Min sch-Q1-Q2 peptide reporting the mean residue ellipticity (MRE) in the 190–250 nm interval. Overlay of spectra obtained at increasing temperatures (5, 25, 50, 75°C) and subtraction (\( \Delta \)) of the spectra measured at 5°C and 75°C, either in saline buffer (two left panels) or in the same buffer with 80% v/v TFE (two right panels).

(E) As in panel D for the Rhi fer-Q1-Q2 peptide.

(F) Plots of the 222 nm MRE over increasing temperatures (5°C–75°C) with (dark gray) or without TFE (light gray) for Min sch-Q1Q2 (left panel) and Rhi fer-Q1Q2 (right panel). MRE was normalized at its value at 5°C.

(G) Graphs plotting the difference (\( \Delta \)) in normalized MRE as a function of temperature, either in the presence or absence of TFE, as calculated from the graphs in panel F, for Min sch-Q1Q2 and Rhi fer-Q1Q2. See also Figure S5.

The previous findings prompted us to define mechanistically the structural and functional impact of the observed evolutionary polyQ length variation in FOXP2. While the FOXP2 DNA-binding domain structure is solved,\(^{36} \) that of the polyQ-bearing N-terminus is unknown. Existing evidence strongly suggests that the FOXP2 polyQ repeats may form \( \alpha \)-helical coiled coils (CCs) whose stability may be length-dependent and reduced by proline residues.\(^{30,36} \)

To determine the possible CC structure of the FOXP2 polyQ repeats and the impact of their variation, we first obtained structural predictions of the CC propensity of human FOXP2 and chiropteran orthologs with representative polyQ variation patterns (Figure 4A), i.e., expansion (Cynopterus), contraction (Miniopterus) and contraction by proline/polyP insertion (Rhinolophus).

Paircoil2\(^{37} \) identified in human FOXP2 four high-probability CC regions, i.e., cc1-4 (with 1 – P-score) \( \geq 0.8 \)\(^{37} \) consistent with \( \alpha \)-helical structure predictions (Figure 4B). Cc2 and cc3, overlap with Q1 and Q2, respectively, whereas cc4 is a known leucine zipper.\(^{37} \) While cc1 and cc4 are conserved across orthologs, the extension of cc2 and cc3 varies with polyQ length (Figures SSA–SSD). Proline residues in Rhinolophus widen the gap between cc2 and cc3, normally caused by a single P within the hinge peptide. Atomic level structural predictions using Raptor-X\(^{92} \) (Figures 4C and S5E) are in agreement with Paircoil2 in showing Q1 and Q2, and flanking Q-rich regions, forming an antiparallel CC hairpin
of two \( \alpha \)-helices separated by a turn (hinge). The Cynopterus ortholog displayed the longest CC with Q-Q hydrogen bonds\(^{93} \) and the proline region in Rhinolophus forms a random coil loop between Q\(_1\) and Q\(_2\).

To experimentally test these predictions, we chemically synthesized Q\(_1\)Q\(_2\) peptides (Figure 4A). Given the challenges in synthesizing/solubilizing long polyQ repeats,\(^{94} \) we started from two peptides with short (Rhi fer) and medium (Min sch) polyQ length. While the first displayed good solubility, Min-sch-Q\(_1\)Q\(_2\) was already much less soluble, producing an opalescent solution indicating the prevalence of oligo-/polymers.\(^{95} \) Based on these observations, we did not further attempt to synthesize longer peptides, and we performed a comparative analysis of Min-sch-Q\(_1\)Q\(_2\) and Rhi-fer-Q\(_1\)Q\(_2\).

The CD spectra of Min-sch-Q\(_1\)Q\(_2\) in saline buffer displayed \( \alpha \)-helical structure with a random coil component (Figure 4D). The helical component was progressively disrupted by heating from 5°C to 75°C, with an isodichroic point at \(~203\) nm, indicative of an equilibrium between \( \alpha \)-helical and random coil conformations.\(^{30} \) The subtraction of 5°C and 75°C spectra shows that the structured component disrupted by heating has CC signatures, i.e., a ratio between the ellipticities at 222 and 208 nm > 1 (e.g., Fiumara et al.\(^{36} \); Figure 4D) indicating that the helical signal derives from a mixture of CC oligomers and random coil protomers rather than by partially helical protomers. Trifluoroethanol (TFE, 80% v/v) stabilized as expected the helical structure, even at 75°C, with a 222/208 nm ellipticity ratio >1, indicative of CC formation (Figure 4D), as also observed in the case of polyserine (polyS).\(^{96} \) A subtraction spectrum under these conditions (Figure 4D) had a single, red-shifted 222 nm peak and a blunted 208 nm peak indicative of CC polymers.\(^{30,95,96} \)

Consistent with predictions, the Rhi-fer-Q\(_1\)Q\(_2\) spectra revealed a lesser helical signal at 5°C, more easily destabilized by heating (Figure 4E). Subtraction spectra displayed a 222/208 nm ellipticity ratio <1, which may suggest the prevalence of \( \alpha \)-helices over CC oligomers, although the random coil signal related to the predicted long, disordered proline-rich loop may also deepen the 208 nm peak, thus masking the CC signal (Figure 4E). This interpretation is corroborated by experiments in TFE in which subtraction spectra displayed a 222/208 nm ellipticity ratio >1 and a blunted 208 nm peak (Figure 4E), indicative of CCs in a polymeric state,\(^{95} \) as above.

The differential stability of the \( \alpha \)-helical structures of the two peptides was also highlighted by the analysis of the absolute ellipticity at 222 nm upon heating (Figures 4F and 4G).

Altogether, these structural analyses show that the FOXP2 polyQ repeats and flanking peptides form \( \alpha \)-helical structures with CC propensity, whose stability and oligomerization are modulated by polyQ length variation and proline insertion patterns as observed throughout phylogeny.

**Atomic force microscopy (AFM) reveals condensates and fibrillar structures formed by FOXP2 polyQ peptides**

To further define the organization of their oligo-/polymers at the morphological level, we studied the Min sch-Q\(_1\)Q\(_2\) and Rhi fer-Q\(_1\)Q\(_2\) peptides by means of AFM (Figure 5).

The AFM analysis of Min-sch-Q\(_1\)Q\(_2\), which forms an opalescent solution, was disturbed by numerous large peptide assemblies distributed unevenly on the mica surface, which made the scanning of smaller particles difficult, either in saline buffer or in the presence of TFE. Figure S5A exemplifies one of these, which may derive from tangling of fibrillar structures as previously found for polyA CC peptides.\(^{36} \) In areas free from large assemblies, we detected spherical droplets interspersed with thin fibrillar profiles (Figure 5A). The frequency of fibrillar profiles appeared to be increased by TFE (Figure 5A). The droplets ranged in diameter from a few nanometers to mesoscale condensates of tens-hundreds of nm (Figure 5B) which have been associated with LLPS.\(^{97} \)

The more soluble Rhi fer-Q\(_1\)Q\(_2\) peptide was easier to analyze. Interestingly, it formed complex structures with tangled fibrils emerging radially from a rounded central condensate, a transitional morphology between globular and fibrillar structures (Figure 5A). Notably, similar structures have been observed in AFM of proteins undergoing LLPS.\(^{103} \) The fibrillar component was enhanced by TFE, with the formation of large bundles of long fibrils (Figure 5A), consistent with the increase in the 222/208 nm ellipticity ratio in CD subtraction spectra measured in the presence of TFE (Figure 4E, right panel).

Together with the CD results, these findings indicate that CC \( \alpha \)-helical peptides containing the FOXP2 polyQ repeats can assemble into LLPS-like mesoscale condensates and elongated fibrils, with transitional forms, as illustrated in Figure 5B. The relative proportion of these structures can be modulated by polyQ length, as indicated by the differential abundance of transitional structures formed by the two peptides. These results are consistent with evidence that CCs can drive the formation of both LLPS-driven condensates\(^{34,35} \) and fibrils,\(^{30,95,96} \) and that polyQ assemblies can transition from liquid-like condensates to more solid fibrillar structures.\(^{105} \)

**Human FOXP2 forms nuclear LLPS-driven condensates**

The previous results, and existing evidence that CCs and amino acid repeats can drive LLPS,\(^{34,35} \) prompted us to test whether FOXP2 can undergo LLPS in the cellular context.

Toward this aim, we first tested whether FOXP2 can form discrete foci in cells, and then whether these foci have recognized, distinctive properties of LLPS-driven condensates (Figure 6).

Using immunocytochemistry and super-resolution confocal microscopy (Airyscan) to reveal the subcellular distribution of FOXP2, we found that the endogenous protein forms numerous discrete foci of variable size in the nuclei of both HEK293 and Neuro-2a cells (Figure 6B, upper panels). These results were confirmed upon expression of GFP-tagged FOXP2, which formed similar foci that became larger and more defined in a time-dependent manner (Figure 6B, lower panel), as typical of proteins undergoing intracellular LLPS.\(^{104} \)

Based on this morphological evidence, we further tested whether the observed FOXP2 nuclear foci exhibit typical features of LLPS-driven condensates.\(^{35,102} \) First, by performing fluorescence recovery after photo bleaching (FRAP) experiments (Figure 6C), we found that the
FOXP2-GFP foci undergo rapid molecular exchange with substantial FRAP in 90–120 s (half-time 16.83 ± 1.92 s) and a mobile fraction of 78%, like other LLPS-prone proteins. Second, we found that the foci are sensitive to 1,6-hexanediol (1,6-HEX; Figure 6D), which disrupts the relatively weak molecular interactions underlying LLPS, as indicated by a ~75% reduction of the relative condensate area in the nuclei of cells treated with 1,6-HEX for 15 min in comparison with the nuclei of control cells (0.24 ± 0.04, n = 24 fields, vs 1 ± 0.06, n = 12, respectively, values normalized to control, p < 0.0001, t-test).

These findings indicate that FOXP2 can form LLPS-driven condensates in the cellular context.

**The polyQ-bearing N-terminal domain of FOXP2 drives LLPS and can recruit FOXP1**

Our CD and AFM in vitro analyses strongly suggest that the polyQ repeats may drive FOXP2 LLPS. To test this hypothesis in the cellular context, we determined whether the isolated polyQ-bearing CC N-terminal domain (NTDCC) can recapitulate the LLPS behavior of FOXP2. When we expressed GFP-tagged NTDCC in cells, we found that it forms rounded foci as full-length FOXP2. The foci were also present

![Figure 5. Morphological analysis of Rhi fer-Q1Q2 and Min sch-Q1Q2 by AFM](image-url)
**Figure 6.** FOXP2 and its polyQ region undergo intracellular LLPS and recruit FOXP1 into condensates

(A) Schematic representation of the constructs for the cellular expression of human FOXP2 protein or fragments of it, in fusion with either GFP or mCh-Cry2. Those that were expressed in the study of intracellular LLPS. Together with a construct for the expression of the full-length protein (fl, 715 a.a.) in fusion with GFP, we generated constructs for the expression of the NTD bearing three CC (ctt-cc-3 regions (NTDcc, a.a. 1-244), the same NTD devoid of the polyQ repeats (ΔQ2Q3) and the polyQ fragment alone (Q2Q3), all in fusion with either GFP or mCh-Cry2.

(B) Upper panels. AnyScan super-resolution confocal fluorescence images of nuclei of a Neuro-2a cell (larger image) and a HEK293 cell (smaller images, including DAPI staining in blue) after immunostaining of endogenous FOXP2. Note how the protein forms discrete foci (arrowheads) of variable size in the nucleus. A few foci in the cytoplasm may derive from a known, alternatively spliced isoform of FOXP2 (‘10+’) devoid of the C-terminus that localizes to the cytoplasm. Lower panels. Confocal fluorescence images of cell nuclei of different HEK293 cells expressing FOXP2-GFP for 24 (left), 48 (middle), and 72 (right) hours. Note how the protein forms similar foci (arrowheads) as in the upper panels that become more defined with increasing time after transfection, as typical of LLPS-prone proteins.

(C) Molecular mobility within FOXP2-GFP foci as detected by FRAP. Left panel: confocal image of the nucleus of a FOXP2-GFP expressing cell in which target regions of interest (ROIs) are outlined by yellow lines. One of the ROIs was photobleached (p) while the other was used as a control (c). The middle panel shows confocal images of the two ROIs taken at the indicated timepoints before (pre) and after photobleaching. Right panel: FRAP curve of FOXP2-GFP.

(D) Left panel: confocal fluorescence images of cells expressing FOXP2-GFP, either treated (+) or not (-; controls) with 1,6-Hex. The arrowhead indicates a condensate in control cells. The arrow indicates fragmented; residual condensates after 1.6-Hex application. Calibration bar: 10 μm. Right panel: bar graph representing the relative area occupied by condensates in nuclei of transfected cells in 1.6-Hex-treated (+) versus control (-) cultures. Data are represented as mean ± SEM, and the statistical tests that were performed are reported in the results section. Asterisks indicate statistically significant differences. The same applies to panels E and G.

(E) As in panel D, but for the NTDcc-GFP construct. The arrowhead and arrow indicate, respectively, a small and a large condensate. Note how larger condensates are more resilient to 1,6-Hex. Calibration bar: 10 μm.

(F) Left panel: time-lapse confocal images of HEK-293 cells expressing NTDcc-mCh-Cry2 (upper panels) or mCh-Cry2 (controls; lower panel) acquired before (pre) and after (1 and 20 min) photoactivation (arrowhead). Right panel: graph plotting the temporal kinetics of condensate formation and dissolution, quantified as the relative cell area occupied by condensates, up to 20 min after photoactivation in cells expressing either NTD-mCh-Cry2 (dark gray) or mCh-Cry2 (light gray). Calibration bar: 10 μm.

(G) Confocal fluorescence microscopy images of cells expressing mCh-Cry2 alone (controls) or NTDcc ΔQ2Q3 or Q2Q3 in fusion with mCh-Cry2, 2 min after photoactivation with 488 nm light (left panel). The bar graph (right panel) plots the relative cell area occupied by condensates in cell cultures expressing the four constructs that were either non-phototoactivated (controls; gray bars) or photoactivated (black bars) with 488 nm light (values normalized to the non-phototoactivated control for each group). Calibration bar: 10 μm.

(H) Left three panels: confocal fluorescence microscopy images of the nucleus of a HEK293 cell co-expressing FOXP2-GFP and FOXP1-DsRed. Note in the overlay image (merge) the high degree of co-localization of the two proteins within the same condensates. (see also Figure S7D). Right three panels: as in the left panels but in cells co-expressing FOXP2 NTDcc-GFP and FOXP1-DsRed. Calibration bar: 10 μm. See also Figure S7.

Figure 6. FOXP2 and its polyQ region undergo intracellular LLPS and recruit FOXP1 into condensates

(A) Schematic representation of the constructs for the cellular expression of human FOXP2 protein or fragments of it, in fusion with either GFP or mCh-Cry2. Those that were expressed in the study of intracellular LLPS. Together with a construct for the expression of the full-length protein (fl, 715 a.a.) in fusion with GFP, we generated constructs for the expression of the NTD bearing three CC (ctt-cc-3 regions (NTDcc, a.a. 1-244), the same NTD devoid of the polyQ repeats (ΔQ2Q3) and the polyQ fragment alone (Q2Q3), all in fusion with either GFP or mCh-Cry2.

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(C) Molecular mobility within FOXP2-GFP foci as detected by FRAP. Left panel: confocal image of the nucleus of a FOXP2-GFP expressing cell in which target regions of interest (ROIs) are outlined by yellow lines. One of the ROIs was photobleached (p) while the other was used as a control (c). The middle panel shows confocal images of the two ROIs taken at the indicated timepoints before (pre) and after photobleaching. Right panel: FRAP curve of FOXP2-GFP.

(D) Left panel: confocal fluorescence images of cells expressing FOXP2-GFP, either treated (+) or not (-; controls) with 1,6-Hex. The arrowhead indicates a condensate in control cells. The arrow indicates fragmented; residual condensates after 1.6-Hex application. Calibration bar: 10 μm. Right panel: bar graph representing the relative area occupied by condensates in nuclei of transfected cells in 1.6-Hex-treated (+) versus control (-) cultures. Data are represented as mean ± SEM, and the statistical tests that were performed are reported in the results section. Asterisks indicate statistically significant differences. The same applies to panels E and G.

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(F) Left panel: time-lapse confocal images of HEK-293 cells expressing NTDcc-mCh-Cry2 (upper panels) or mCh-Cry2 (controls; lower panel) acquired before (pre) and after (1 and 20 min) photoactivation (arrowhead). Right panel: graph plotting the temporal kinetics of condensate formation and dissolution, quantified as the relative cell area occupied by condensates, up to 20 min after photoactivation in cells expressing either NTD-mCh-Cry2 (dark gray) or mCh-Cry2 (light gray). Calibration bar: 10 μm.

(G) Confocal fluorescence microscopy images of cells expressing mCh-Cry2 alone (controls) or NTDcc ΔQ2Q3 or Q2Q3 in fusion with mCh-Cry2, 2 min after photoactivation with 488 nm light (left panel). The bar graph (right panel) plots the relative cell area occupied by condensates in cell cultures expressing the four constructs that were either non-phototoactivated (controls; gray bars) or photoactivated (black bars) with 488 nm light (values normalized to the non-phototoactivated control for each group). Calibration bar: 10 μm.

(H) Left three panels: confocal fluorescence microscopy images of the nucleus of a HEK293 cell co-expressing FOXP2-GFP and FOXP1-DsRed. Note in the overlay image (merge) the high degree of co-localization of the two proteins within the same condensates. (see also Figure S7D). Right three panels: as in the left panels but in cells co-expressing FOXP2 NTDcc-GFP and FOXP1-DsRed. Calibration bar: 10 μm. See also Figure S7.

To determine the contribution of the polyQ repeats to the NTDcc-driven LLPS, we generated NTDcc deletion constructs (Figure 6A) for the expression of the polyQ repeat region alone (Q2Q3-mCh-Cry2), or of the NTDcc devoid of it (ΔQ2Q3-mCh-Cry2). We compared the relative capacity of these two fragments to induce LLPS with that of the whole NTDcc by measuring condensate formation in cultures 2 min after photoactivation relative to untreated control cultures (Figure 6G). Overall, a two-way ANOVA revealed significant effects related to the expressed construct, photoactivation, and their interaction (F1,105 = 45.50, F3,105 = 180.66, F2,213 = 45.50, respectively, p < 0.001 in all instances) in the induction of condensates upon photoactivation in comparison with basal levels.

Notably, LLPS induction was considerably lower for Q2Q3-mCh-Cry2 in comparison with the NTDcc, showing that the polyQ repeats have an important role in driving FOXP2 LLPS (4.25 ± 0.33, vs. 10.61 ± 0.86 at 2 min, n = 20 fields, values normalized to the mean pre-activation relative condensate area in each group, p < 0.001, NK test). Indeed, the polyQ region alone (Q2Q3-mCh-Cry2) was still able to drive significant LLPS induction (4.46 ± 0.64 at 2’ post-activation vs. 1.00 ± 0.15 pre-activation, n = 10 in each group, values normalized to pre-activation levels, p < 0.001, NK test), although at lower levels in comparison with the whole NTDcc (4.46 ± 0.64, n = 10, vs. 10.61 ± 0.86 at 2 min, n = 20, values
transfected cell area in comparison with high (comparisons; Figure 7C). The mean condensate size also varied similarly across constructs (Figure 7C; F(2, 10321) = 222.34, p < 0.001). Similar FOXP1 recruitment into condensates even in the cytoplasm (Figures 6H and S7D).

Results where obtained when only the NTD CC domains of the three constructs were expressed (F(2,86) = 25.19, p < 0.001, one-way ANOVA; for human FOXP2 and FOXP1. We found that the two proteins can co-condensate in cell nuclei and that the NTC CC alone was able to mediate cooperation with them in driving LLPS.

Photoactivation induced a less extensive LLPS of G giants were expressed as full-length proteins or as NTD CC fragments, in fusion with either GFP or mCh-Cry2 (Figure 7A).

Size). Consistent with these findings, the three NTD CC variants displayed differential ability to undergo triggered LLPS (Figure 7D). Indeed, FRAP experiments further revealed that the mobility of molecules within condensates was higher for the RHs variant (F(2,45) = 23.93, p < 0.001, repeated measures ANOVA; Figure 7F), with a significantly different at 1 post-bleaching between the two constructs (p < 0.03; NK test). Moreover, the RH FRAP curve displayed, in comparison with the HS one, a significantly lower time constant normalized to pre-activation levels in each group, p < 0.001, NK test). Thus, also the Q-rich regions flanking the Q1Q2 repeats (Figure 1A) cooperate with them in driving LLPS.

As functionally related proteins can undergo LLPS and co-condensate within the same compartments,35 we tested whether this is the case for human FOXP2 and FOXP1. We found that the two proteins can co-condensate in cell nuclei and that the NTC CC alone was able to mediate FOXP1 recruitment into condensates even in the cytoplasm (Figures 6H and S7D).

These findings identify the NTD CC and its polyQ repeats as drivers of FOXP2 LLPS and highlight their role in the recruitment of interactors into condensates, consistent with the role of polyQ CCs in mediating homo-/heterotypic protein interactions.30,36

**Evolutionary polyQ length variation regulates FOXP2 LLPS**

The previous evolutionary analyses strongly suggested that polyQ variation may regulate not only FOXP2 structure but also its function in terms of LLPS and transcriptional activity. To test this hypothesis, we generated constructs for the expression of chimeric variants of human FOXP2 (Hs) in which the polyQ repeats were either replaced with those of other species representative of the observed polyQ variation patterns, i.e., *Cynopterus sphinx* (Cs), *Miniopterus schreibersii* (Ms), or *Rhinolophus ferrumequinum* (Rf), or deleted (ΔQ1ΔQ2 variant). These variants were expressed as full-length proteins or as NTD CC fragments, in fusion with either GFP or mCh-Cry2 (Figure 7A).

We first compared the ability to form condensates of the GFP-tagged human FOXP2 (Hs) with that of evolutionary variants with low (Rf) or high (Cs) polyQ length (Figure 7B). We found that the Rf and Cs condensates occupied, respectively, a lower and higher percentage of the transfected cell area in comparison with Hs condensates (F(2,86) = 23.10, p < 0.001, one-way ANOVA; mean relative cell area occupied by condensates: 1 ± 0.04, n = 33 fields, for HS; 1.41 ± 0.08 for CS, n = 12; 0.78 ± 0.04 for RF, n = 46; p < 0.02, for both the Cs vs. HS and RF vs. HS comparisons; Figure 7C). The mean condensate size also varied similarly across constructs (Figure 7C; F(2, 10321) = 222.34, p < 0.001). Similar results were obtained when only the NTD CC domains of the three constructs were expressed (F(2,86) = 25.19, p < 0.001, one-way ANOVA; mean relative cell area occupied by condensates: 1 ± 0.06, n = 32, for HS, 1.55 ± 0.10 for CS, n = 26, 0.75 ± 0.07 for RF, n = 31, p < 0.03, for both the Cs vs. HS and RF vs. HS comparisons, NK post-hoc test; F(2,430) = 32.408, p < 0.0001, one-way ANOVA, for the mean condensate size). Consistent with these findings, the three NTD CC variants displayed differential ability to undergo triggered LLPS (Figure 7D). Indeed, photoactivation induced a less extensive LLPS of RHs-NTD CC-mCh-Cry2 compared to HS-NTD CC-mCh-Cry2, while the opposite was true for CS-NTD CC-mCh-Cry2. Indeed, a two-way ANOVA (Figure 7E) revealed significant differences (p < 0.001) related to the construct (F(3,158) = 75.62), photoactivation (F(1,158) = 478.19), and their interaction (F(3,158) = 75.62). Moreover, significant differences (p < 0.001; NK test) were found between all constructs and the control (mCh-Cry2), showing that all variants displayed significant LLPS induction upon photoactivation, and between HS and either CS or RF.

Moreover, FRAP experiments further revealed that the mobility of molecules within condensates was higher for the RF than for HS variant (F(2,45) = 23.93, p < 0.001, repeated measures ANOVA; Figure 7F), with a significant difference at 1 post-bleaching between the two constructs (p < 0.03; NK test). Moreover, the RF FRAP curve displayed, in comparison with the HS one, a significantly lower time constant.
Evolutionary polyQ variation regulates FOXP2 transcriptional activity

Finally, we determined the possible regulatory effect of the evolutionary polyQ variation on the transcriptional activity of FOXP2. FOXP2 regulates several target genes, including SRPX2, another speech-/language-associated gene, whose expression is repressed by FOXP2. Indeed, FOXP2 has at least six binding sites in the SRPX2 promoter (Figure 7G).

Thus, we used luciferase reporter assays to measure the relative transcriptional activity of the FOXP2 polyQ variants (i.e., Hs, Rf, Ms, Cs, and ΔQ1Q2) as in Roll et al. HEK293 cells were co-transfected with a pcDNA4/HisMax expression vector encoding for any one of the FOXP2 polyQ variants, together with a second vector encoding for firefly luciferase (Luc−) under the control of the SRPX2 promoter (SRPX2-pGL3-luc), and a third vector for the CMV promoter-driven expression of Renilla luciferase (RLuc; pRL-CMV plasmid) as a transfection control. The empty pcDNA4/HisMax vector was transfected to measure basal luciferase levels in control cultures (Figure 7G).

Then, we compared the relative luciferase luminescence in cultures expressing FOXP2 polyQ variants. A one-way ANOVA indicated overall significant differences between the different polyQ variants (F(5,338) = 29.10). Post hoc comparisons confirmed, as expected based on previous observations, that Hs-FOXP2 induces a significant reduction of Luc− expression in comparison with control cultures, consistent with the repressive role of FOXP2 on SRPX2 (mean relative luminescence 0.79 ± 0.03, n = 69 culture wells, vs. 1 ± 0.02, n = 116, values normalized to the mean control value, p < 0.001 NK post hoc test; Figure 7H).

Notably, Rf and ΔQ1Q2, with low or null polyQ length, displayed a significantly increased repressive activity in comparison with all other variants bearing longer polyQ tracts, including Hs-FOXP2 (p < 0.01 in all instances, NK test; Figure 7H). Overall, we observed a correlation between PolyQ length and transcriptional repression (r = −0.97, n = 5, p < 0.01; Figure S7E). In species with predominant Q1 length variation (Rf and Ms vs. Hs), SRPX2 repression also correlated with the Q1/Q2 ratio (Figure S7F).

These findings indicate that the evolutionary polyQ variation may encode vocalization frequency at the molecular level by regulating quantitatively the transcriptional activity of FOXP2.

DISCUSSION

This study identifies a quantitative relationship between molecular and vocalization-related phenotypic parameters throughout mammalian evolution by which the FOXP2 Q1/Q2 length ratio encodes vocalization frequency, from the infrasonic to the ultrasonic ranges. Consistent with this finding, we observed generalized patterns of convergent polyQ length evolution related to vocalization/hearing frequency in FOXP2 and, for certain taxa, in FOXP1. At the molecular level, we found that the polyQ region of FOXP2 forms α-helical CCs driving LLPS. Finally, we observed that polyQ length variation impacts both CC structure and LLPS, ultimately regulating the transcriptional activity of FOXP2. These findings open new perspectives on the molecular underpinnings of vocalization and its evolution.

Molecular encoding of vocalization frequency in the length of polyQ repeats in FOXP2 and FOXP1

Our analyses indicate that the frequency range of laryngeal vocalization in mammals is quantitatively encoded by FOXP2 polyQ lengths, so that lower Q1/Q2 length ratios indicate higher frequencies, and vice versa.

The correlation that we found between FOXP2 polyQ lengths and laryngeal vocalization frequency, or related morpho-functional parameters, was overall considerably robust and generalized in Chiroptera and, more broadly, in Mammalia, with some outliers at the species or taxon level. In this latter respect, two taxa that appeared to be outliers in this overall correlation either use non-laryngeal forms of USVs, i.e., Odontoceti, or display a parallel system of frequency encoding in the polyQ repeats of FOXP1, i.e., Muridae. The latter form of encoding based on the FOXP1 polyQ repeats is prevalent in Rodentia and Afrotheria, and is limited to a few species in other mammalian taxa.

Some previous studies have shown qualitatively how the species-specific occurrence of amino acid repeats can be related to interspecific divergence, but have not analyzed the effect of their length variation across species. Other studies have instead identified quantitative correlations of polyQ length variants with morphological or physiological parameters but only among individuals of the same species. Our findings extend these analyses by systematically studying the length variation of amino acid repeats in a large number of species, identifying an overall quantitative correlation between polyQ lengths and vocalization frequency, and extensive patterns of polyQ convergent evolution across mammalian taxa.

Convergent evolution of polyQ repeats

Our evolutionary analyses identify extensive patterns of polyQ length convergence at the molecular level that correspond, at the organismal level, to convergence in vocalization/hearing frequency ranges across mammals.

In molecular evolution studies, convergence is generally defined in terms of single amino acid substitutions which can only be associated qualitatively, as a binary feature, to the presence of a certain phenotype, such as ultrasonic vocalization. Conversely, in our analyses, polyQ length variation could be quantitatively related overall to vocalization/hearing frequency ranges. At the same time, the polyQ sum-vs-ratio
plots also captured qualitatively the nuanced vocalization features of Chiroptera (NE, FM1, FM2, CF). Therefore, this analytical approach offers powerful quantitative and qualitative tools in molecular evolution studies.

A growing body of evidence indicates that polyQ repeats are enriched in neural proteins, displaying complex evolutionary dynamics throughout phylogeny.16,40,41,111 The evidence of convergent evolution adds a novel element to our appreciation of the complex evolutionary history of polyQ repeats.

The patterns of polyQ convergence between species in relation to vocalization/hearing frequency can be detected within and between mammalian orders. This is already evident in Chiroptera, in which we analyzed >50 species, eight of which display phenotypic convergence in USV type with species not belonging to their same superfamily. In six of them, this is paralleled by polyQ convergence in FOXP2, while in Pteronotus parnellii and Noctilio leporinus we observed polyQ convergence with USV-emitting rodents in FOXP1. P. parnellii (Noctilionoidea), which emits CF USVs, also has a unique Q-to-P substitution extending the predicted loop between Q1 and Q2, as in CF-emitting Rhinolophoida. Strikingly, similar polyQ convergence patterns in FOXP2 and/or FOXP1 were found between species in most mammalian orders. Thus, these findings identify a first case of extensive molecular convergence associated with vocalization-related organismal phenotypes.

In mammals, multiple cases of molecular convergence of single residue substitutions are known in hearing-related proteins (e.g., prestin) in taxa with high-frequency vocalization/hearing, i.e., echolocating Chiroptera and Cetacea.48–52,112 However, none of the convergent mutations in these individual proteins identifies all the echolocating mammals in other taxa.51,52 Our findings highlight widespread polyQ convergence related to vocalization/hearing frequency across mammals, over the entire infrasonic to ultrasonic frequency range.

In contrast to hearing-related proteins, evidence of convergent molecular evolution in vocalization-related proteins is limited. Besides our findings, there is evidence of convergence between Chiroptera and Cetacea in muscle proteins involved in USV emission.53 For FOXP2, convergence between echolocating Chiroptera and Cetacea has not been observed,28 which may be related to the distinct USV emission modalities used by species belonging to these two taxa. Indeed, Chiroptera generate laryngeal USVs emitted through the mouth or nose, whereas more stereotyped (ultrasonic clicks in Odontoceti are generated by non-laryngeal phonic lips and emitted through the frontal lipid-filled melon.28 The latter route does not permit rapid orofacial and sensorimotor coordination, processes in which FOXP2 is implicated.12,113 Thus, USV emission in Cetacea may require molecular adaptations in genes other than FOXP2, which underwent accelerated evolution in Chiroptera.28 Our findings, in showing no polyQ convergence between Chiroptera and Cetacea (Odontoceti), are thus consistent with this view and with the notion that FOXP2 is more closely linked to the evolution of laryngeal vocalization.

Interestingly, polyQ/polyA variation in RUNX2 has been related to cranial morphological parameters in certain orders, like Carnivora and Primates,39,114,115 but not in others,116,117 indicating taxon-specific roles of the RUNX2 repeats. The patterns of FOXP2 polyQ convergence instead extend across mammals at large, although some degree of taxonomic specificity emerged, as for some USV-emitting Rodentia. Similarly, substantial variation in FOXP1 polyQ repeats is generalized in Rodentia and Afromerodermia but restricted to fewer species in other taxa (Chiroptera, Lagomorpha, Eulipotyphla). Interestingly, some species display USV-related polyQ changes in both FOXP2 and FOXP1.

The case of the pika (O. princeps) highlighted the predictive power of FOXP2 polyQ lengths for vocalization/hearing frequency ranges. Indeed, its polyQ repeats converge with those of USV-emitting bats, in striking parallelism with the known pika-bat similarity in the evolution of the hearing-related protein prestin.72 We found that wild O. princeps emit calls extending into the ultrasonic range and has morphological cochlear parameters indicative of high-frequency hearing.72 These findings rationalize the pika-bat convergence in both FOXP2 and prestin.

Coiled coils, LLPS, transcription, and their polyQ length-dependent regulation

To understand mechanistically how the evolutionary changes of the polyQ stretches may affect the FOXP2 protein, we first analyzed the structural and functional features of these repetitive sequences, and then defined the impact of their length variation at the molecular level.

We found that the two major polyQ repeats and the interposed hinge peptide form α-helical CCs with length-dependent stability. Indeed, CD analyses of two polyQ peptide variants showed that both exist in an equilibrium between random coil, α-helical, and CC conformations, that can be shifted by polyQ length variation, in agreement with existing evidence on polyQ sequences.36,39,53,96,118,119 These results, consistent with atomic-level structural predictions, support a model in which the two polyQ repeats form α-helices separated by a loop (hinge peptide) assembling into an antiparallel CC hairpin. The extension of the loop, and thus CC stability, can be varied by interspersed proline residues, which appeared recurrently throughout phylogeny in Q1 or, in the unique case, within the hinge peptide of P. parnellii which contains two proline residues instead of one as in other species. Remarkably, besides the mere polyQ expansion/contraction, Q-to-P substitutions and polyP insertions, are virtually the only evolutionary changes that we observed in the FOXP2 polyQ repeats. This can be rationalized by considering that proline destabilizes α-helices and CC-driven oligomerization.90,120,121 Proline insertions in Q1 may thus represent a recurrent molecular adaptation regulating CC length, stability, and oligo-/polymerization.

Overall, variable combinations of Q1 and Q2 length changes lead to a graded variation of their total and relative lengths. Such polyQ CC length variation may represent a structural rearrangement ultimately regulating protein function,32,33,36,108 and a notion that is supported by our findings.

The AFM morphological analysis of the assemblies formed by FOXP2 polyQ peptides revealed both rounded mesoscale condensates and long fibrillar structures with transitional figures, as found in AFM studies of LLPS-prone proteins.98 Remarkably, the relative proportion of condensates and fibrils is modulated by polyQ length and by a helix-stabilizing agent (TFE). In CD spectra, TFE enhanced the α-helix/CC signal, which displayed signatures associated with fibrillar CC polymers. These findings together indicate that α-helix/CC polymers is closely related to the oligo-/polymerization modalities of FOXP2 polyQ peptides. Thus, while relatively unstable α-helices/CCs may drive LLPS through less regular interactions between protomers (e.g., CC assembly by helix swapping),109 more stable α-helices/CCs may promote
fibrillization through more regular structural arrangements (e.g., CC oligomer stacking). This model is consistent with the observation of transitional profiles between condensates and fibrils and with the notion that LLPS may represent a nucleation step triggering fibrillation.

The AFM findings of condensate-like structures were confirmed in the cellular environment. Indeed, both endogenous and exogenously expressed FOXP2 form discrete, 1,6-HEX-sensitive intranuclear foci which undergo rapid FRAP, as typical of LLPS-driven condensates. The polyQ-bearing NTD (and its Q1+Q2 fragment) recapitulated the phase separation behavior of the entire protein. Moreover, the NTD alone was also able to recruit FOXP1 into condensates. These findings concurrently identified the NTD, and its polyQ repeats as the drivers of FOXP2 LLPS. Remarkably, human FOXP2 mutants in which the polyQ repeat lengths were changed to mimic their evolutionary variation displayed differential ability to undergo LLPS as a function of total and relative polyQ lengths.

These findings are consistent with the emerging ability of polyQ and other amino acid repeats in transcription factors to drive LLPS, as shown for the polyQ/polyA repeats of RUNX2. Notably, we had found that also these repeats in RUNX2 form z-helical CCs, with length-dependent stability, that regulate protein oligo-/polymerization and transcriptional activity. Thus, the evolutionary length variation modulating CC stability may be a general mechanism to regulate transcription factor activity.

This interpretation is supported by our observation that, besides LLPS, the FOXP2 transcriptional activity is also related to the total and relative polyQ lengths. Indeed, lower Q1+Q2 and Q1/Q2 were associated with higher repressive activity on the target SRPX2 promoter and vice versa. Only two studies have previously analyzed the possible regulatory roles on transcription of the FOXP2 polyQ repeats with conflicting results. These studies focused only on repeats deletions, one of which is very modest (i.e., a single Q deletion), and did not test a series of polyQ length variants. Estruch et al. reported that polyQ repeats do not significantly regulate transcription, despite some variable effects that were observed. However, in this study the luciferase experiments were performed using FOXP2 variants with N-terminal YFP tags, which may interfere by steric hindrance with the NTD structure and function. This may explain the small, variable transcriptional effects observed upon polyQ deletion. Conversely, Zhao et al. proposed that a single Q deletion may abolish the repressive effect of FOXP2 on the CNTNAP2 promoter. However, the same authors reported very different expression levels of the FOXP2 variants in their cell lines, which correlated with CNTNAP2 levels, putting into question their conclusion. Thus, a single Q deletion would have no effect when normalizing for expression levels (see also Estruch et al.). Possible cell type- and promoter-specific transcriptional effects of FOXP2 should also be taken into account in comparing the results of the different studies.

Remarkably, our results are instead similar to those obtained in experiments studying a FOXP1 variant devoid of its polyQ repeats, which displayed enhanced repressive activity. Moreover, a recent comparison of the transcriptional activity of multiple polyQ variants in the androgen receptor (AR) highlighted length-dependent transcriptional effects. These and our findings suggest the possibility that the more LLPS-prone polyQ variants of FOXP2 may be more easily retained into condensates and less able to bind target promoters, thus limiting their regulatory effect on transcription, although alternative mechanisms cannot be ruled out.

Toward a quantitative understanding of the molecular underpinnings of mammalian vocalization

A growing body of evidence indicates that polyQ and other amino acid repeats can modulate in a length-dependent manner, the structure, interactions, and function of proteins. Consistent with these views, the results of our analyses provide evidence that the observed evolutionary polyQ length variation regulates the structure, LLPS, and transcriptional activity of FOXP2.

Taken together, our findings uncover a quantitative molecular encoding of vocalization frequency in the relative lengths of polyQ repeats in FOXP2, highlight widespread patterns of vocalization-related polyQ convergent evolution, and rationalize mechanistically how polyQ length variation can have phenotypic impact by regulating the molecular function of FOXP2.

These results and analytical approaches contribute to a quantitative, mechanistic understanding of vocalization and its evolution.

Limitations of the study

Our analyses of the molecular evolution of FOXP2 and FOXP1 polyQ repeats in relation to vocalization frequency-related parameters were deliberately focused on Mammalia, given their considerable degree of polyQ length variability and their highly diversified vocalization modalities and frequency ranges. Thus, the study did not include some non-mammalian taxa, such as Aves, in which a considerable number of species are known to emit highly sophisticated and diversified forms of vocalization. We are currently exploring whether the results of our analyses can also apply to these non-mammalian taxa.

STAR+METHODS

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Supplemental information can be found online at https://doi.org/10.1016/j.isci.2023.108036.

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AUTHOR CONTRIBUTIONS
Conceptualization, design, and supervision of the study: F.F.; Data collection and experimental work: S.V., V.V., M.D., C.M., F.C., D.M., L.L., I.P., and F.F.; Data analysis and interpretation: S.V., V.V., M.D., C.M., F.C., D.M., L.L., I.P., F.R., F.J.M., L.C., M.G., and F.F.; Manuscript writing: F.F. with input from all authors; Funding acquisition: F.F. All authors approved the submitted version. S.V., V.V., and M.D. contributed equally to this work and are co-first authors. Corresponding author and lead contact: F.F. (ferdinando.fiumara@unito.it).

DECLARATION OF INTERESTS
The authors declare no competing interests.

INCLUSION AND DIVERSITY
We support inclusive, diverse, and equitable conduct of research.

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REFERENCES


## STAR METHODS

### KEY RESOURCES TABLE

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## Experimental models: Cell lines

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### Software and algorithms

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RESOURCE AVAILABILITY

Lead contact
Further requests for information, resources, and reagents should be directed to and will be fulfilled by the lead contact, Ferdinando Fiumara (ferdinando.fiumara@unito.it).

Materials availability
Reagents generated as part of this study, as listed in the key resources table, are available from the lead contact upon request.

Data and code availability
- The analyses presented in Figures 1, 2, 3, and S1–S4 are based on i) publicly available protein and nucleotide sequences, obtained from the NCBI (www.ncbi.nlm.nih.gov) and Uniprot (www.uniprot.org) databases, ii) animal vocalization frequency and body mass datasets as derived from the cited literature, and iii) µCT cranial scans, obtained from the Morphosource database (www.morphosource.org), either as freely accessible data or upon authorization from dataset administrators. The accession numbers of FOXP2 and FOXP1 protein or nucleotide sequences are listed in Table S1. The vocalization frequency and body mass datasets were derived from the literature, as listed (with PMID identifiers) in Tables S2 and S3. The MorphoSource specimen list is reported in Table S4. The data related to structural analyses, microscopy and other experiments reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to re-analyze the data reported in this paper is available from the lead contact upon request.
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cells
HEK293 cells (293-F strain; Thermo Fisher) and Neuro-2A cells (a kind gift of G. Merlo, University of Turin) were maintained following standard procedures at 37°C with 5% CO2 in DMEM (Thermo Fisher) supplemented with 10% fetal bovine serum, 2 mM L-glutamine, 100 units/mL penicillin, 100 μg/mL streptomycin, and, for Neuro-2A cells, MEM non-Essential amino acids solution (1X; Gibco).

METHOD DETAILS

Bioinformatics
PolyQ lengths were quantified using a polyQ analyzer Perl script in protein sequences of FOXP1/2 orthologs from Uniprot (www.uniprot.org) or NCBI (www.ncbi.nlm.nih.gov/protein/). For certain species, polyQ lengths were manually determined by translating genomic sequences identified through BLAST searches on the NCBI genome platform (https://www.ncbi.nlm.nih.gov/genome/) using human FOXP2 or FOXP1 sequences as the query. The list of species for which FOXP1/2 polyQ repeats were analyzed, their name abbreviations, and the corresponding protein or genomic sequence IDs are listed in Table S1. For each FOXP1/2 ortholog, we calculated the length of the two polyQ repeats (‘Q1’ and ‘Q2’) corresponding in sequence alignments to the two major repeats flanking the hinge peptide ‘ALQVARQLLL’ in human FOXP1 or ‘HPQKQAKE’ in human FOXP2. In a few predicted FOXP2 sequences of Cetacea species, the hinge peptide is extended by a likely spurious 22-residue peptide (‘VGSGRLTHAEEGEAGRGPRRPG’ in T. truncatus, with minor substitutions in other species), which is not present in other cetacean or mammalian orthologs. It is encoded by a genomic sequence immediately downstream of a conserved splice site located at the end of the sequence encoding the canonical hinge octapeptide. This splice site may have been missed in the computational identification of FOXP2 exon-intron junctions in certain Cetacea genomes. Sequencing of FOXP2 cDNAs from Cetacea will be required to address this uncertainty in a conclusive manner. Anyhow, the presence or absence of this peptide does not change Q1 and Q2 lengths in these predicted ortholog sequences, which were therefore included in our analyses.

For each ortholog, we calculated the Q1+Q2 sum and the Q1/Q2 ratio. In the case of multiple available sequences/isoforms for each species, we analyzed the sequence with the longest polyQ tracts. The Q1 repeats of FOXP2 are interrupted by one or more proline residues in several species. In these species, given the structural effect of proline residues, we calculated the Q1 length as the length of the uninterrupted polyQ repeat upstream of the first proline residue, while fragmented repeats between this proline residue and the hinge peptide were not taken into account. In a few cases, a single histidine residue was occasionally found within Q1. Given the polar nature of the residue, it was considered as being part of the polar polyQ stretch rather than an interruption of it.

Sequence alignments were generated using MultAlin. Protein domain schemes were obtained from MyDomains and modified using Photoshop Elements 11 (Adobe). Phylogenetic trees were derived from TimeTree and elaborated using Mega X. Silhouette drawings in the public domain of species of interest were downloaded and modified from PhyloPic (http://phylopix.org). M. ungulatus and O. degus silhouettes were derived from Wikimedia (https://commons.wikimedia.org; pictures by Pacos and Algesirdas).

Phenotypic datasets
Vocalization modalities, vocalization frequency/bandwidth, and body mass for Chiroptera were derived from Collen. The relative cochlear size (CW/BW in Figure 1D) for chiropteran species was derived from Simmons et al. The maximum audible frequency (60 dB SPL) in audio-sequences identified through BLAST searches on the NCBI genome platform (https://www.ncbi.nlm.nih.gov/genome/) using human FOXP2 as the query. The list of species for which FOXP1/2 polyQ repeats were analyzed, their name abbreviations, and the corresponding protein or genomic sequence IDs are listed in Table S1. For each FOXP1/2 ortholog, we calculated the length of the two polyQ repeats (‘Q1’ and ‘Q2’) corresponding in sequence alignments to the two major repeats flanking the hinge peptide ‘ALQVARQLLL’ in human FOXP1 or ‘HPQKQAKE’ in human FOXP2. In a few predicted FOXP2 sequences of Cetacea species, the hinge peptide is extended by a likely spurious 22-residue peptide (‘VGSGRLTHAEEGEAGRGPRRPG’ in T. truncatus, with minor substitutions in other species), which is not present in other cetacean or mammalian orthologs. It is encoded by a genomic sequence immediately downstream of a conserved splice site located at the end of the sequence encoding the canonical hinge octapeptide. This splice site may have been missed in the computational identification of FOXP2 exon-intron junctions in certain Cetacea genomes. Sequencing of FOXP2 cDNAs from Cetacea will be required to address this uncertainty in a conclusive manner. Anyhow, the presence or absence of this peptide does not change Q1 and Q2 lengths in these predicted ortholog sequences, which were therefore included in our analyses.

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Morphological analyses on µCT scan analyses

Micro computed tomography (µCT) scans encompassing the cranial region of species of interest were obtained from the MorphoSource database (https://www.morphosource.org; Media IDs: 86463,96992,2150,170220,170399,170595,165459,02723,158814,13131,11011,10956,168375,15460,15477,15612,15720,410414,394893,140436,8620,114319,56554,22387,16145,16139,16131,13626,22292,22235,21964,21967,21967,21952,45544,167374,55626,55624,47005,167378,55698,79025,367903,47052,55618,55349,167384,365356,365356,167937,36511,36128,170595,57668,59128; Table S4; The P. alecto scan S9128 was used only to generate the left panel in Figure 1C). For each scan, the image stack was cropped to the region of interest, converted into 8-bit format, and saved in Nifti format using ImageJ.141 The cropped scans were used to generate digital endocasts of the cochlea and lateral semicircular canals using ITK-SNAP after segmentation based on manual thresholding. Cursors were positioned on landmarks of interest and 3D coordinates were used to calculate distances between them, and their ratios, using Excel. For each scan, we measured the cochlear height-to-width ratio18 and the cochlear width/basicranium width (CW/BW) ratio.18 For most Primates/Lagomorpha species, mean values of these parameters were obtained from 2 or more specimens for each species. In the case of Leporidae, the mean values were derived from one specimen each of Lepus europaeus, L. articus, L. americanus, and correlated with available polyQ parameters derived from L. europaeus. For Ochotonidae, the mean values were derived from three specimens of Ochotona princeps and one of O. pallasi and correlated with the available polyQ parameters from O. princeps. For Chiroptera, the morphological parameters were obtained from a single specimen of each species of interest (see Table S4).

Vocalization detection and analysis

To analyze spontaneous vocalizations of wild American pikas (Ochotona princeps), we analyzed archived recordings originally taken as part of a monitoring project (Cascades Pika Watch; https://scistarter.org/cascades-pika-watch) of a resident population at talus slopes in the Columbia River Gorge (OR, USA). Pika vocalizations were recorded using an acoustic microphone with some sensitivity to ultrasonic signals (up to 25 kHz) connected to a SM3BAT recorder (Wildlife Acoustics). The recordings that were analyzed, showing evidence of pika vocalizations extending in the ultrasonic range with harmonics >20 kHz (see Figure 2D), were recorded between May 29 and June 5, 2017 (databasin.org, “2017 Western Columbia River Gorge” dataset), using a microphone located near Herman Creek (microphone coordinates: 45° 40’ 24.0384”, 121° 50’ 15.2884”). The vocalization spectrograms were analyzed using the Kaleidoscope software (Wildlife Acoustics).

Structural predictions

Predictions of FOXP2 secondary structure were obtained using DeepCNC143 on the Quick2D platform (https://toolkit.tuebingen.mpg.de/). Predictions of coiled coil structures were obtained using Paircoil2.90 Atomic-level structural models for a.a. 1–240 FOXP2 orthologs were chemically synthesized (Thermo Fisher Scientific) with N-terminal acetylation, C-terminal amidation and purity >95%. Peptides were dissolved in a saline buffer (100 mM NaCl, 10 mM phosphate buffer, pH 7.4) to generate stock solutions (1 mg/mL) which, after the circular dichroism measurements, were aliquoted, flash-frozen in liquid N2, and stored at −80°C. The Miniopterus-derived peptide formed an opalescent solution whose concentration was difficult to measure spectrophotometrically. Thus, peptide concentrations in circular dichroism experiments were estimated based on peptide weight. This may overestimate the concentration of soluble, optically active chiral material and thus underestimate the ellipticity signal.19 Before each measurement, peptides were further diluted in saline buffer to a working concentration of 0.2 mg/mL, transferred to quartz cuvettes (1-mm optical length), and CD spectra were collected in the 190–260 nm frequency range (every 0.5–1 nm) using a J-815 spectropolarimeter (Jasco). In some measurements peptides were diluted in saline buffer containing 50–80% v/v 2,2,2-trifluoroethanol (TFE, Sigma-Aldrich). Blank spectra of saline buffer, or of saline buffer with TFE, were subtracted from the peptide spectra in the same buffers. The mean residue molar ellipticity [θ] was calculated as [θ] = θ × mw/(10 × (n–1) × c × pL), where θ is the measured ellipticity, mw is the molecular weight of the peptide, n is the number of amino acids in the peptide, c is the concentration of the peptide (mg/mL), and pL is the cuvette pathlength (cm).146 To test structural stability, the samples were heated (from 5°C to 75°C; 10°C/min) while recording ellipticity at 222 nm. Data and graphs were elaborated using Spectra Analysis (Jasco) and Excel (Microsoft).

Atomic force microscopy (AFM)

AFM scans were performed under ambient conditions in tapping mode by using a sharp Al-coated silicon cantilever (Tap190 AL-G, Budget Sensors; length: 225 μm; width: 38 μm; tip radius: ca. 10 nm, 10° at the apex), near the resonance frequency (190 kHz), mounted on an Easyscan2 AFM (Nanosurf) equipped with a high resolution 10-μm scan head, within a shielded and insulated enclosure placed on an antivibration platform, as in Pelassa et al.36 Peptide stocks in saline buffer, stored at −80°C, were thawed and diluted to 0.05–0.25 mg/mL in saline buffer either with or without 50–80% TFE. The diluted solutions were incubated at 4°C for 10–15 min and a few microliters of them were dropped onto freshly cleaved mica (V1 grade muscovite, Ted Pella) and let air dry for ~5 min. The mica surface was gently rinsed with ultrapure water and dried under a mild stream of nitrogen. Samples were analyzed as soon as the surface became dry, within 1 h. In some measurements
Addgene (#101221).

For all plasmids that were generated, the coding sequences of FOXP2 and tags werecompetent cells and purified using the QIAprep Spin Mini-/Maxi-prep kits (Qiagen), or the NucleoSpin plasmid Mini kit (Macherey-Nagel), QIAquick (Qiagen) or the NucleoSpin (Macherey-Nagel) kits. Plasmids were amplified by transformation of Stellar (Clontech) chemically

The pcDNA4/HisMax-FOXP2 plasmid encoding human FOXP2 and the pGL3-SRPX2 containing the SPRX2 promoter were kindly provided by Dr. P. Szeptetowski (INSERM, France). The pcDNA4/HisMax-FOXP1 plasmid encoding human FOXP1 was kindly provided by Dr. A. Banham (University of Oxford, UK). The pRL-CMV vector was obtained from Promega (#e2261). The pHR-mCh-Cry2WT plasmid was obtained from Addgene (#101221).

Plasmids encoding chimeric polyQ variants of human FOXP2, in which the sequence encoding Q₁, Q₂, and the hinge peptide were replaced by the corresponding sequence of chiropteran FOXP2 orthologs of interest, were obtained using the In-Fusion cloning system (Clontech-Takara). Briefly, two PCR products with overlapping ends (15 bp overlaps) were obtained using the PrimeSTAR-GXL-SP DNA Polymerase (Clontech). The first product consisted of the pcDNA4/HisMax-FOXP2 plasmid sequence devoid of the region encoding the two major polyQ repeats (residues 152–209 in the human protein). The second product contained a fragment of FOXP2 orthologs of species of interest (i.e., C. sphinx (Cs), M. schreibersii (Ms), or R. ferrumequinum (Rf)) overlapping with the human FOXP2 polyQ region fragment (a.a. 152–209) in FOXP2 ortholog sequence alignments.

For the Cs and Ms orthologs, these products were PCR-amplified using long, partially overlapping forward and reverse primers, encoding overall for the entire region of interest (as derived from NCBI). For the Rf ortholog, the second product was PCR-amplified from a plasmid encoding the region of interest previously obtained by gene synthesis (GeneArt, Thermo Fisher). All the three primer pairs used for the Cs, Ms, and Rf polyQ region contained 15 bp extensions, corresponding to the ends of the pcDNA4/HisMax-FOXP2-derived PCR product, suitable for cloning using the In-Fusion system. Finally, the latter PCR product was alternatively combined with one of the Cs, Ms or Rf polyQ fragments in cloning procedures using the In-Fusion system to generate the pcDNA4/HisMax-FOXP2-Cs, -Ms, and -Rf plasmids. These plasmids were used, together with the original plasmid encoding the Homo sapiens (Hs) ortholog, that we called pcDNA4/HisMax-FOXP2-Hs. In a fifth construct (ΔQ₁Q₂), the polyQ region (a.a. 152–216, i.e., from the start of Q₁ to a few residues after the end of Q₂) was deleted by re-circularizing a pcDNA4/HisMax-FOXP2-derived PCR product upon ligation using a T4 ligase (NEB). These five plasmids (i.e., pcDNA4/HisMax-FOXP2-Hs, -Cs, -Ms, -Rf, -ΔQ₁Q₂) were then used for luciferase assay experiments. Moreover, they served as templates for the PCR amplification of sequences encoding full-length human FOXP2 polyQ variants (FOXP2-Hs, -Cs, -Ms, -Rf) with suitable 15 bp extensions for their In-Fusion cloning into the pcAcGFP1-N In-Fusion Ready vector (Clontech-Takara; #632501) for the expression of the GFP-tagged polyQ variant proteins. These constructs were used for studying the subcellular distribution of FOXP2 polyQ variants (confocal fluorescence microscopy) and their transcriptional activity (luciferase assays). A set of plasmid constructs was similarly generated, using pAcGFP-N1-In-Fusion-Ready as the recipient plasmid, for the expression of GFP-tagged N-terminal domains (NTDs) of human FOXP2 and its polyQ length variants (Cs, Ms, Rf and ΔQ₁Q₂), corresponding to the first 244 residues of human FOXP2 in ortholog protein alignments, which encompass the first three CC regions in the Paircoil2 prediction (NTDcc constructs; Figures 4A and 7A).

We generated a plasmid for the expression of FOXP1-DsRed by PCR-amplifying the FOXP1 coding region from the pcDNA4/HisMax-FOXP1 plasmid with suitable extensions for In-Fusion-cloning into the pDsRed-monomer-N1 In-Fusion Ready (Clontech-Takara, #632498).

To use the optoDroplet system for a spatiotemporally controlled induction of protein LPLS in living cells, we generated a vector for the cellular expression of proteins of interest fused to mCherry (mCh; for visualization) and to an A. thaliana Cry2 fragment (Cry2; for photoactivation). The DNA sequence coding for the mCh-Cry2 fusion was PCR-amplified from the pHHR-mCh-Cry2WT plasmid (Addgene #101221), with 15 bp extensions suitable for In-Fusion cloning into a pDsRed-monomer-N1 vector (Clontech-Takara), from which we had previously enzymatically excised the dsRed coding sequence using BamHI and NotI (NEB). The resulting pX-mCh-Cry2 vector was linearized by digestion with XhoI (NEB) and used as a recipient vector for PCR fragments of interest bearing suitable 15 bp extensions for In-Fusion cloning and encoding protein fragments of interest to be expressed as fusions with mCh-Cry2. Using this approach, we generated constructs for the expression of the human FOXP2 NTDcc fragment, its sub-fragments (i.e., Q₁Q₂ (a.a. 146–221) and ΔQ₁Q₂), and its polyQ variants (Cs, Ms, Rf) as described above.

All PCR products in the cloning procedures were separated by agarose gel electrophoresis and bands of interest were purified using QiAquick (Qiagen) or the NucleoSpin Spin (Macherey-Nagel) kits. Plasmids were amplified by transformation of Stellar (Clontech) chemically competent cells and purified using the QiAprep Spin Mini-/Maxi-prep kits (Qiagen), or the NucleoSpin plasmid Mini kit (Macherey-Nagel), or NucleoBond Xtra MIDI EF kit (Macherey-Nagel). For all plasmids that were generated, the coding sequences of FOXP2 and tags were verified by Sanger sequencing.

Cell culture transfection and fixation

For immunocytochemistry and imaging of fluorescently tagged proteins, cells were plated on poly-L-lysine-coated (0.1 mg/mL) glass coverslips within 24-well culture plates. For optoDroplet and FRAP experiments, cells were plated into poly-L-lysine-coated dishes obtained starting from Falcon 50 mm plastic cell culture dishes (Corning) that were perforated to produce a central hole (10 mm) then closed by a glass coverslip glued using Sylgard 184 (Dow Corning). For luciferase reporter assay, HEK293 cells were cultured in 96-well white plates (VWR).
transfection was performed in non-adherent cells in suspension immediately before plating using the same reagents. Two-three days after transfection, the cell cultures were either used for live-cell imaging (FRAP, optoDroplet induction) or fixed. Before fixation, cell cultures on coverslips were rinsed with phosphate-buffered saline (PBS, pH 7.4), fixed with 4% paraformaldehyde (ThermoFisher) in PBS for 15 min at room temperature (RT), rinsed with PBS (3X) and mounted on slides using the Dako fluorescent mounting medium (Agilent).

**Immunocytochemistry**

HEK293 and Neuro-2a cells, fixed 24–48 h after plating, were permeabilized (0.1% Triton X- in PBS, 15 min), rinsed in PBS (3X), and incubated in blocking solution (2% bovine serum albumin (BSA; Sigma) in PBS (1 h at RT). After rinsing in PBS (3X), the cells were incubated with a primary anti-FOXP2 antibody (ThermoFisher, #720031; 1:100 in 0.1% BSA in PBS) for 1 h at RT, rinsed in PBS (3X), and then incubated with fluorescently labeled (Alexa 488) anti-rabbit secondary antibodies for 1 h at RT. After rinsing in PBS (3X), coverslips were mounted on slides. In some experiments, coverslips were also incubated with DAPI (Sigma) for 15 min at room temperature and then rinsed with PBS (3X) before mounting.

**Confocal fluorescence imaging and fluorescence recovery after photo bleaching (FRAP)**

Confocal fluorescence imaging was performed using Olympus FV300 or Leica TCS SP5 confocal microscopes. Super-resolution imaging analyses were performed using a ZEISS LSM 800 Airyscan microscope system using default Airyscan settings. In general, each experiment was replicated at least three times and images were collected from multiple coverslips per experiment.

FRAP experiments were performed using the TCS SP5 microscope, and the FRAP Wizard module of the LAS-X software (Leica). Fluorescent foci formed by GFP-tagged FOXP2 polyQ variants in HEK293 cell nuclei (48–72 h after transfection) were brought into the focal plane, laser power and acquisition parameters were adjusted to avoid signal saturation, and one of the foci was included in a region of interest (ROI) to be photobleached. After acquiring three pre-bleaching images of the cell nucleus, one per second, the ROI was photobleached using the 488 nm laser (100% power) for 3 s, and fluorescence recovery was monitored by acquiring images of the nucleus every second for 120 s. After image acquisition, fluorescence levels in both photobleached and 2–3 control areas of each nucleus were quantified using the LAS-X software and normalized to their respective pre-bleaching fluorescence levels (i.e., the mean fluorescence level in the 3 pre-bleaching scans). To correct for slow photobleaching upon repeated laser scanning during the experiment, the values of the photobleached area were further normalized, at each time point, to the mean value of the non-bleached control areas. In some experiments, the mobility of foci and nuclei on the z axis caused major distortions of the typical fluorescence recovery profiles and/or sudden disappearance of the foci from the ROI. These measurements, for both the Hs and Rf constructs, were not included in the analysis. Data were analyzed using Easy-FRAP for curve fitting, and r-half and immobile fraction calculations.

**Liquid-liquid phase separation (LLPS) analysis**

The spontaneous formation of condensates by fluorescently tagged human FOXP2, its fragments, and polyQ variants, was quantified by confocal fluorescence imaging in HEK293 cells 48–72 h after transfection. Maximum intensity projections of z stack fluorescence confocal images of microscopy fields (233x233, or 350 x 350 μm) were converted into 8-bit images using ImageJ (NIH). Image brightness and contrast were adjusted to highlight either the foci or the entire nuclei/cell profiles of the fluorescent cells. These images were used to automatically quantify, for each field, the surface area occupied by condensates and that occupied by nuclei/cells using ad hoc CellProfiler pipelines. The ratio between the two areas defined the proportion of the cell area occupied by condensates.

To define the sensitivity to 1,6-hexanediol (1,6-HEX) of the FOXP2-GFP and FOXP2-NTD-GFP foci, which is typical of LLPS-driven condensates, we used the protocol by Ullanov et al. Briefly, HEK293 cell cultures (48 h after transfection) were transiently permeabilized with 1% Tween 20 and exposed to 5% 1,6-HEX for 10 min. Control cultures were treated in the same manner, except that 1,6-HEX was omitted. The cells were then rinsed with PBS and fixed in 4% PFA in PBS (15 min at RT). The coverslips were mounted onto microscope slides and, after confocal imaging, the area occupied by foci was measured using CellProfiler, as described above.

The triggered induction of LLPS of protein fragments of interest was obtained using the optoDroplet system. Briefly, cells were transfected with plasmids suitable for the expression of FOXP2 fragments of interest fused to mCh-Cry2, as described above, and LLPS was induced 48–72 h after transfection by 488 nm light illumination using two distinct protocols. To quantify the overall LLPS induction in cell populations, the cell cultures were photoactivated 48–72 h after transfection using blue light (488 nm), illuminating the entire coverslip, emitted by the 100 W fluorescent lamp of an Eclipse TE200 microscope (Nikon) with suitable dichroic filters for 5–7 min. Two minutes after photoactivation, the cells were rinsed in PBS and fixed for subsequent fluorescence confocal imaging to detect the mCherry red fluorescence. Control cells were treated in the same way, except that photoactivation was omitted. LLPS-driven condensate formation was quantified, as described above, by calculating for each construct the relative area of nuclei (or full-length FOXP2 or its polyQ variants), or cells (for FOXP2 NTDccc fragments), occupied by condensates in photoactivated cultures compared to non-photoactivated control cultures.

To quantify the kinetics of LLPS induction, reversibility, and re-induction, we performed live-cell imaging experiments using an Olympus FV300 confocal microscope. In these experiments, the culture medium was replaced with HBSS, pH 7.4 (Sigma) at RT and the red mCh fluorescence of 2–10 cells within a magnified confocal microscopy field (at an intermediate z level of the cell layer) was monitored using 561 nm laser light, every minute before (5 scans) and after (20 scans) a 60 s photoactivating pulse delivered by fast-scanning the sample with blue light (488 nm; 20% laser power). In some experiments, to test LLPS reinduction, a similar pulse was delivered 20 min after the first one and red fluorescence was monitored again for 20 min. LLPS-driven condensate formation was quantified, as described above, by calculating the
relative area of nuclei (for full-length FOXP2 or its polyQ variants), or cells (for FOXP2 N-terminal fragments), occupied by condensates in cultures before and after photoactivation. For each nucleus/cell, all values were normalized to the mean value in the scans before the first photo-activating pulse.

**Luciferase transcription reporter assay**

For luciferase transcription reporter assays, HEK293 cells were plated into 96-well white culture plates (VWR). Twenty-four hours after plating, the cells were co-transfected with three plasmid vectors, i.e., (i) the pcDNA4/HisMax plasmid, either empty (control condition) or encoding human FOXP2 (Hs), or one of its polyQ variants (Cs, Ms, Rf, and ΔQ1Q3); (ii) the pGL3-SPRX2 plasmid; (iii) the pRL-CMV vector (Promega) as a transfection control. For transfecting 8 culture wells, we mixed 1125 μg of the empty pcDNA4/HisMax plasmid (or equimolar amounts of the same vector encoding human FOXP2 or its polyQ variants), 255 μg of pGL3-SPRX2, and 120 μg of pRL-CMV. The DNA mix was transfected in a 3:1 M:ml ratio with Fugene-6 or Lipofectamine 2000, following the manufacturers’ protocols. Forty-eight hours after transfection, firefly and Renilla luciferase activity were sequentially measured using the Dual-Glo luciferase assay system (e2929; Promega) and a GloMax microplate reader (Promega). For each well, the firefly luciferase luminescence was first normalized against the Renilla luciferase luminescence to correct for transfection efficiency. Then, all luminescence values were further normalized to the average value measured in the experimental control group expressing the empty pcDNA4/HisMax plasmid. In some luciferase assays, GFP-tagged FOXP2 variants were expressed and the pEFGP-C1 vector, instead of pcDNA4/HisMax, was transfected in control cultures. At least 18 wells from three independent experiments were analyzed for each FOXP2 variant construct. Each experiment included control wells for normalization (pcDNA4/HisMax control), for a total of 116 control wells across all the luciferase assay experiments.

**Software**

Basicranium μ-CT stacks shown in Figure 1C were obtained using the Aleph 3D-viewer embedded in the MorphoSource website (www.morphosource.org). Ochotona princeps vocalization recordings were analyzed using Kaleidoscope Lite 5.4.7 software (Wildlife Acoustics). Digital endocasts of the left cochlea and lateral semicircular canals, as shown in Figures 2E and 2F, and landmark coordinates were obtained using ITK-SNAP 3.8.0. Gwyddion 2.57 was used to analyze AFM experiments. UCSF Chimera 1.43 was used to analyze, and generate images of protein structure models generated by Raptor-X. Circular dichroism data were analyzed using Spectra Manager (version 2.06.01; JASCO Corporation) and Excel (Microsoft). Confocal fluorescence microscopy images were visualized and processed using ImageJ (NIH), Fluoview (Olympus), LAS-X (Leica) and ZEN lite (Zeiss). ImageJ and Photoshop Elements 11 (Adobe) were used for image processing and to generate figures.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Data are expressed as mean ± standard error of mean (SEM). The details concerning the specific parameters that were quantified in each analysis (e.g., polyQ lengths, vocalization-related and morphological parameters, condensate area, luminescence), the number of experimental groups, their composition (e.g., species, cells, condensates, culture wells) and numerosity (n), are reported above, where appropriate, in the results and method details sections. Student’s t test, one- or two-way ANOVA, ANOVA for repeated measures, Newman-Keuls post hoc test, and other tests, were performed where appropriate, as detailed in the results section. In all instances, a p value ≤ 0.05 was considered statistically significant. Data analysis and statistics were performed using Excel (Microsoft), Statistica (Tibco) and Python 3 (pandas, matplotlib, and numpy packages).

**Accession numbers**

The accession numbers of protein or nucleotide (genomic) sequences of the FOXP2 and FOXP1 orthologs that were analyzed are reported in Table S1. Micro computed tomography (μCT) scans were obtained from the MorphoSource database (https://www.morphosource.org). The scan Media IDs (also listed in Table S4) are: 86463, 96992, 2150, 170220, 170399, 170595, 165459, 02723, 158814, 13131, 11011, 10956, 168375, 15460, 15477, 15612, 15720, 410414, 394893, 140436, 8620, 114319, 56554, 22387, 16145, 16139, 16131, 13626, 22292, 22235, 21964, 21967, 21967, 21952, 45544, 167374, 55626, 55624, 47005, 379025, 379023, 47052, 55618, 55349, 167384, 365356, 167937, 36587, 57664, 167934, 36439, 36171, 25920, 25943, 36828, 167937, 36257, 36434, 36511, 36128, 76668, 59128, The P. alecto scan S9128 was used only to generate Figure 1C, left panel.