Neurofibromatosis type 1 (NF1; OMIM 162200) is one of the most common neurogenetic conditions, affecting 1 in 3,000 people worldwide. Characterized by a propensity to develop nervous system tumors, learning and behavioral deficits, and pigmentary abnormalities, NF1 is caused by a germline sequence alteration in the NF1 gene (OMIM: 613113; chromosome 17q11.2). In addition, somatic NF1 sequence changes have been reported in numerous other cancers, extending their importance to malignancies occurring in the general population.

The NF1 gene, comprising 57 exons, was initially identified, and its complete coding sequence was assembled over 25 years ago. In the ensuing decades, 4 alternatively spliced exons (9a, 10a-2, 23a, and 48a) were discovered and added to a large number of exons with temporary numerical assignments (e.g., 10a2 or 23.2). This has led to considerable confusion and inconsistencies in clinical and scientific publications, which also has critical implications for NF1 DNA sequence alteration reporting as part of patient management, genotype-phenotype correlations, and the interpretation of small sequence changes. To provide a unified annotation system for the NF1 gene, the mouse and human genes were aligned and assembled using published sequences for sequential numbering and comparison.

The full-length cDNA sequences of NF1 (8457 nucleotides; figure e-1 at Neurology.org/ng) and NF1 (8463 nucleotides; figure e-2) genes were initially aligned using Nucleotide Basic Local Alignment Search Tool (BLAST) (NIH) optimized for megablast, revealing 92% sequence identity with 0.08% mismatches (figure e-2). Excluding the alternatively spliced exons, the ubiquitously expressed 57 exons correspondingly encode 2818 amino acids, while the corresponding full-length mouse Nfi transcript, Nfi-003 (ENSMUST000000108251.8), comprises 57 exons and encodes 2,820 amino acids, with 2 additional amino acids encoded by exon 17. Alignment of the predicted amino acid sequences of the human (RefSeq NM_000267; UniProt P21359) and mouse (UniProt Q04690) full-length transcripts using protein BLAST revealed 98% amino acid sequence identity (figure e-3).

Next, we examined the previously published alternatively spliced exons: exon 9a contains a 30-nucleotide sequence encoding 10 amino acids and is predominantly expressed in postmitotic brain neurons, whereas the muscle-specific exon 48a contains 54 nucleotides that encode 18 amino acids at the carboxyl terminus of the protein. The 63-nucleotide exon 23a encodes 21 amino acids that are inserted within the RAS-GTPase-activating protein (GAP)-related domain of the NF1 protein (neurofibromin), resulting in diminished RAS-GAP activity. The most recently described alternatively spliced exon, exon 10a-2, contains 45 nucleotides and encodes 15 amino acids, is located at the amino terminus of neurofibromin, and this sequence has been hypothesized to direct intracellular membrane targeting by virtue of its predicted transmembrane domain. Using published alternatively spliced exon sequences (figure e-1), BLAST searches were performed, and these alternatively spliced exons were renumbered and named according to their location (figure 1)—between exons 11–12 (11alt12; formerly 9a), exons 12–13 (12alt13; formerly 10a-2), exons 30–31 (30alt31; formerly 23a), and exons 56–57 (56alt57; formerly 48a). In this regard, we propose that the nucleotides (and the encoded amino acids) of these alternatively spliced exons be numbered 11alt12_1–30 (1–10), 12alt13_1–45 (1–15), 30alt31_1–63 (1–21), and 56alt57_1–54 (1–18). Possible new alternatively spliced exons should be similarly named based on their location.

11alt12 is 100% identical to both the cDNA and amino acid sequences of the predicted Mus musculus transcript variant X7 (mRNA XM_006532442.3), while 30alt31 is 98.4% identical to the cDNA and 95.2% identical to the amino acid sequence (containing a single conservative lysine-to-arginine change) of Mus musculus Nfi (ENSMUST00000071325.8) (figures e-4 and e-5). Notably, the 12alt13 and 56alt57 human sequences were not found in either mouse transcriptome or genome databases following
megablast, discontinuous megablast, or BLASTN searches (figures e-4 and e-5). Based on the absence of 12alt13 and 56alt57 sequences in the mouse, further studies will require to define their roles in neurofibromin function. It should be noted that neither 12alt13 nor 56alt57 has been reported in other reference model organisms (rat or zebrafish; data not shown). Taken together, we assembled and updated the human NF1 and mouse Nf1 sequences to facilitate greater consistency in both clinical reporting and basic research studies on NF1. The clarification of the nomenclature for the NF1 gene is a necessary step to enable standardization of exon, nucleotide, and amino acid numbering, which is particularly important for reporting patient-specific DNA sequence variants. In addition, implementation of this proposed numbering system will provide better calibration of human genetic testing results with Nf1 preclinical precision medicine models developed using genetically engineered mice and induced pluripotent stem cells.7

A) Schematic diagram illustrating the 57 exons of the human (top panel) and mouse (bottom panel) full-length transcripts, as well as the positions of the alternatively spliced exons (11alt12, 12alt13, 30alt31, and 56alt57). The proposed numbering system is denoted in each exon, while the prior numbering system is indicated below each exon. The RAS-GTPase-activating protein and the CRAL-TRIO domains are highlighted in gray. (B) Table detailing the nucleotides and amino acids spanning each of the 57 exons in human NF1 and mouse Nf1 genes. CRAL-TRIO domain = cellular retinaldehyde-binding protein and TRIO guanine exchange factor structural domain.

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Author contributions: C.A. performed the analyses, assembled the figures and tables, and helped write the manuscript. L.Q.L. and R.A.K. were involved in the design of the study and edited the manuscript. D.H.G. wrote the manuscript with C.A. and oversaw the study at all stages.

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