INTRODUCTION

Fibrodysplasia Ossificans Progressiva (FOP; OMIM #135100) is an extremely rare autosomal dominant disease in which heterotopic bone forms in muscle and soft tissue, leading to joint dysfunction and severe disability. The incidence of FOP is 1 of 2 million and there is no regional, racial, or sexual predisposition (Baujat et al., 2017; Qi et al., 2017).

Classical presentation of FOP is characterized by congenital malformation of the great toes and progressive heterotopic ossification of soft tissues leading to cumulative disability. The genetic cause of FOP are mutations in the ACVR1 gene that encodes a type I receptor of Bone Morphogenetic Proteins. The most recurrent mutation in FOP patients is R206H affecting the Glycine-Serine rich domain and causing the hyper-activation of the receptor and the responsivity to the non-canonical ligand, Activin A.

In the present study, we described a 3-years old child with early and highly suggestive clinical features of FOP who was found negative for the recurrent p.R206H substitution.

Methods: Molecular screening of the whole ACVR1 coding sequence and functional characterization in transfection-based assays.

Results and Conclusions: We identified a novel, de novo variant in the fifth ACVR1 coding exon (NM_001111067.4:c.772A>T; NP_001104537.1:p.(R258W)). This substitution, never reported in association with FOP, affects a conserved arginine residue in the kinase domain of the protein. In silico analysis predicted the pathogenicity of this substitution, demonstrated by in vitro assays showing that the p.R258W ACVR1 mutated receptor acquires the ability to transduce the aberrant Activin A-mediated signaling, as observed for the gene variants associated with FOP.

KEYWORDS
Activin A, ACVR1, BMP signaling, Fibrodysplasia Ossificans Progressiva, p.R258W

1 | INTRODUCTION

Fibrodysplasia Ossificans Progressiva (FOP; OMIM #135100) is an extremely rare autosomal dominant disease in which heterotopic bone forms in muscle and soft tissue, leading to joint dysfunction and severe disability. The incidence

Serena Cappato and Rasa Traberg contributed equally to this work.
swelling, followed by progressive and disabling heterotopic endochondral ossification (Kaliya-Perumal et al., 2020; Qi et al., 2017). Flare-ups and heterotopic ossification usually manifest in the first decade of life in 95% cases (Baujat et al., 2017). Acute phases caused by the underlying inflammation in the ligaments, tendons, or skeletal muscle occur upon pro-inflammatory insults such as muscle fatigue, tissue damage, intramuscular injections, or viral illness or apparently without any recognizable trigger. They are generally sporadic and unpredictable, in terms of occurrence, duration, and outcome: not all the flare-ups end up with bone neoformation, on the other hand a spontaneous, creeping ossification in absence of acute phases is also observed (Pignolo et al., 2016). Over time, owing to the repeated flare-ups at different sites, progressive and cumulative ossification of soft tissues occurs, leading to the debilitating effects of FOP (Kaliya-Perumal et al., 2020). Some patients have atypical FOP signs, or so-called FOP plus forms: tibial osteochondromas, spinal malformations and broad femoral neck, thumb malformations, cognitive impairment, and diffuse scalp thinning (Kaplan et al., 2009; Qi et al., 2017).

In 2006 the R206H mutation in the ACVR1 (OMIM #102576, also known as ALK2) gene encoding a bone morphogenetic protein (BMP) type I receptor was described as the molecular cause of FOP (Shore et al., 2006). The most common R206H substitution affects the intracellular glycine-serine (GS) domain of the ACVR1 receptor. Atypical missense mutations (p.L196P, p.R202I, p.Q207E, p.R258G/S, p.G328R/W/E, p.G356D, and p.R375P) in the GS or protein kinase domains of ACVR1 have been also identified in some FOP patients (Kaliya-Perumal et al., 2020). De novo heterozygous mutations are found in most cases, but there are familial cases with autosomal inheritance reported. Usually, FOP diagnosis is made on a clinical basis by the observation of the presence of pathognomonic features such as typical toe deformation and ossification foci and is confirmed by DNA sequence analysis of the ACVR1 gene.

2 | MATERIALS AND METHODS

2.1 | ACVR1 screening

Peripheral blood samples were collected upon informed consent administration. Genomic DNA was extracted using the QIAamp DNA Mini kit (Qiagen) according to the manufacturer’s instructions and quantified with the Nanodrop 2000 (Thermo Scientific).

Genomic DNA fragments corresponding to the nine coding exons and flanking intronic sequences of ACVR1 (RefSeq NM_001111067.4; RefSeq NP_001104537.1), were amplified by polymerase chain reaction (PCR; the primer sequences are listed in Table S1) using the GoTaq Master mix (Promega) with reactions carried out in a GeneAmp PCR System 2720 Thermocycler (Applied Biosystems).

PCR products were cleaned up by Exo/SAP-IT (Thermofisher Scientific) digestion and then used for direct sequencing. Reactions were set up with a Big Dye Terminator Cycle Sequencing Kit according to the provided protocol, run on a 3130xl Genetic Analyzer (Applied Biosystems) and sequences analyzed by the Sequencer 4.7 software. Sequence variations were described according to the Human Genome Variation Society guidelines (http://www.hgvs.org).

2.2 | In silico analysis

In order to estimate the impact of the identified variant on ACVR1 receptor function, we performed an in silico analysis by using the Ensembl Variant Effect Predictor (VEP) toolset (available at https://grch37ensembl.org/Tools/VEP) [McLaren et al., 2016].

2.3 | Expression plasmids

Complementary DNA (cDNA) for human, wild-type ACVR1 (Table S1 for oligonucleotides used to generate the product) was obtained by RT-PCR and then subcloned into the expression vector pCMV-3Tag-8 (pCMV_ACVR1) without the native STOP codon, to be in frame with the 3xFLAG tag provided by the vector.

ACVR1 variants p.R206H, p.R258S, and p.R258W were introduced by mutagenesis, by using the QuikChange Lightning Site-Directed Mutagenesis kit (Agilent) according to the provided protocol. All the obtained clones were checked by Sanger sequencing. The generation of the pGL4.17 vector containing the BMP-Responsive Element (BRE) upstream the Luciferase reporter gene (indicated as pGL4.17-BRE) was previously described by our group (Cappato et al., 2016).

2.4 | Cell culture, transient transfection, and treatments

U2OS cells were already available originally obtained from ATCC (HTB-96). Cells were routinely cultured in a complete medium consisting of Dulbecco’s Modified Eagle’s Medium (DMEM), containing 10% fetal bovine serum (FBS, Gibco). Depletion medium was prepared by reducing FBS at 0.5% or by replacing it with 0.1% BSA (Bovine Serum Albumin, Sigma-Aldrich, Merk). Cells were maintained at 37°C in a humidified atmosphere with 5% CO₂.

Transient transfections were performed in 96-well plates, by seeding 5 x 10⁴/well U2OS cells directly in the transfection mix, composed by 30 ng of the pGL4.17-BRE, 40 ng
of pCMV_ACVR1 constructs and 2.5 ng of pGL4.73[hRLuc/SV40] to normalize toward transfection efficiency, using the Lipofectamine 2000 reagent protocol (Invitrogen, Life Technologies). After 4 hr, cells were washed and incubated for additional 16 hr in a fresh medium with 0.5% FBS, and 33 nM ActA or 50 ng/ml BMP2. Next day cells were lysed and processed to evaluate the Luciferase activities with the Dual Luciferase Reporter Assay (Promega) according to the manufacturer's instruction.

**ID1** gene expression and P-SMAD1/5/9 proteins levels were evaluated by seeding 3 × 10^5 U2OS in 6-well plates for transfection with 2 µg of pCMV_ACVR1 constructs using the Lipofectamine 2000 reagent (Invitrogen, Life Technologies). After 48 hr, cells were incubated over-night in a depletion medium containing 0.5% FBS, then activated in a medium containing 0.1% BSA for 3h with 33nM Activin A or 50 ng/ml BMP2 to evaluate **ID1** gene expression and 45 min with 33 nM ActA or 100 ng/ml BMP2 to verify the P-SMAD1/5/9 phosphorylation state.

### 2.5 RNA extraction and quantitative PCR (qPCR)

For expression studies, treated and untreated cells were harvested and total RNA was isolated by using the RNeasy Mini Kit (Qiagen), according to the provided protocol. RNA was quantified with Nanodrop Spectrophotometer (Thermo Scientific), and first-strand cDNA was synthesized by the iScript cDNA Synthesis Kit (BioRad) from 500 ng of total RNA.

Expression of endogenous **ID1** gene was evaluated through qPCR using specific Syber Green strategy (see Table S1 for Oligo sequences, Tib Molbiol Srl). Samples were measured in duplicate and the results were normalized on the geometric mean of **B2M** and **HPRT** genes. qPCR was run on the IQ5 instrument from BioRad and data analysis was performed using the ΔCt method: ratio reference/target = 2^{ΔCt} (Cappato et al., 2019; Livak & Schmittgen, 2001).

### 2.6 Western blot

For detection of the P-SMAD1/5/9 protein level, cells were washed twice with PBS and lysed in 1xRIPA buffer (50 mM Tris HCl pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 1% Sodium Deoxycholic, 0.1% SDS), containing phosphatase and protease inhibitors (PhosSTOP cocktail and Complete tablets, Roche), NaF 50 mM and Na3VO4 1 mM. Protein concentration was determined by the Pierce™ BCA Protein Assay Kit (Thermo Scientific) according to the manufacturer's protocol and 15 µg of total lysates run onto precasted 4%–15% Midi Criterion TGX-gels 18W (BioRad). Proteins were transferred onto Nitrocellulose membrane (BioRad) and probed with the indicated primary antibody at 4°C overnight. After incubation with HRP-conjugated secondary antibodies, protein bands were revealed by chemiluminescence with the Amersham ECL Detection Reagents (GE Healthcare) and detected with the Uvitec instrument (Cambridge, UK). Densitometric analysis of the western blot signals was performed by using the Uvitec software. Primary antibodies were diluted as follows: P-SMAD1/5/9 1:2000 (#13820, SMAD1 1:2000 #9743, Cell Signaling); anti-FLAG M2 monoclonal antibody 1:3000 (#F1804, Sigma-Aldrich, Merk); anti-GAPDH 1:20000 (#MAB374, Millipore).

### 2.7 Statistical analysis

Luciferase reporter gene assays were performed independently three times in triplicate (n = 9). Experiments to evaluate gene expression by qPCR were performed in duplicate from three independent RNA extractions. The analysis of P-SMAD proteins was carried out on three independent experiments. The Unpair t test was applied to verify statistical significance of the observed variations (https://www.graphpad.com/quickcalcs/ttest1.cfm). Significant differences were given as p < .05*, p < .01**, p < .001***.

### 3 RESULTS AND DISCUSSION

#### 3.1 Patient description

The proband is a 3-year-old Lithuanian boy referred to our examination by the orthopedic surgeon. He is the first child of a non-related Lithuanian couple. The pregnancy was uneventful until the 33rd week of gestation when preeclampsia for the mother was diagnosed. The patient was born after emergency Cesarean section. Birth weight was 1580 g (fifth percentile). He was admitted to the newborn intensive care unit due to respiratory distress. Newborn period was otherwise normal.

Soon after birth, hallux deformation of the feet and mild plagiocephaly were noticed. Conventional cytogenetic analysis was performed and revealed a normal male karyotype 46, XY.

The patient was referred to orthopedic surgeon at 4 months of age (2 months of corrected age) and congenital hip dysplasia was diagnosed. Treatment with the Frejka pillow started and continued for 3 months. The X-ray of the feet showed bilateral hallux valgus with accessory rudimentary phalanges (Figure 1a and b). His growth and development were normal and appropriate for age and prematurity. The first lumps appeared at 7 months of age when non-painful flexible masses were noticed bottom to
the lambdoid suture on the occiput. The lump formation progressed, and further examinations were performed at 11 months of age. There were hard palpable masses on the occiput with limited motions of the head. Neurologic examination was normal. Ultrasound of the masses showed enlarged muscle and thickened periosteum. MRI of the head region revealed infiltration of fascia in the site of the lumps (fasciitis; Figure 1c). Endocrinology workup showed no pathologic changes. The next follow-up of the patient was done at 20 months of age and para-spinal, hard subcutaneous lesions of the back were noticed (Figure 1d). The motion of the neck was limited. No new lumps appeared in 1-year period after the last examination.

3.2 Identification and characterization of a novel variant of the ACVR1 gene

The clinical presentation of the proband was highly suggestive of FOP, a rare genetic condition associated with mutations of the ACVR1 gene (Figure 2a). As a first approach, molecular analysis was focused on the search of the recurrent ACVR1 p.R206H substitution with a negative result. Therefore, molecular screening was extended to the whole coding sequence of the gene with identification of the heterozygous c.772A>T (NM_001111067.4) variant in the fifth coding exon, and consequent change of the Arginine 258 by a Tryptophan (NP_001104537.1: p.(R258W); Figure 2b). The mutation was absent in the healthy parents suggesting a likely de novo origin. This variant is not reported in genomic databases and has never been described in association with FOP. However, it affects a highly conserved residue of the tyrosine kinase domain of the protein (Bocciardi et al., 2009). Noteworthy, other disease variants affecting the p.R258 have been already reported: p.R258S (Bocciardi et al., 2009; Eresen Yazıcıoğlu et al., 2013; Morales-Piga et al., 2012; Ratbi et al., 2010) and p.R258G (Kaplan et al., 2015) associated with rare cases of FOP; p.R258G and p.R258M were found in tumor tissues (somatic mutations) from Diffuse Intrinsic Pontine Gliomas (DIPG), Brainstem gliomas, midline astrocytomas, and stomach adenocarcinoma (Buczkwicz et al., 2014; Chang et al., 2016; Taylor et al., 2014).

The change of a positively charged, hydrophilic residue such the Arginine by a Tryptophan with a large aromatic side chain is likely to impact the receptor function. A deleterious impact is also predicted through the sequence analysis performed by the Ensemble Variant Effect Predictor toolset which provides access to a wide collection of genomic annotations and tools suitable to predict the effect of a genetic variant (McLaren et al., 2016). A summary of this analysis is shown in Table 1, where the effect predicted for the novel substitution is compared to that observed for the other R258 variants associated with FOP.

ACVR1 mutation associated with FOP causes an aberrant SMAD1/5/9 signaling and, most importantly, confers to the mutated receptor the ability to respond to a non-canonical ligand, the Activin A (ActA), thus triggering the ectopic bone formation (Hatsell et al., 2015; Hino et al., 2015). As BMPs, ActA belongs to the TGFβ family and is secreted by several cell types including immune cells (Kaplan et al., 2016).

In physiological conditions, ActA does not show osteogenic properties, can bind wild-type ACVR1 without
transducing any intra-cellular signaling, rather negatively affecting the BMP cascade (Hatsell et al., 2015; Lees-Shepard et al., 2018). On the contrary, the mutated receptor triggers the SMAD1/5/9 cascade upon ActA binding (Hatsell et al., 2015; Hino et al., 2015). The acquisition of this neo-function has been demonstrated for the recurrent R206H mutation as well as for other substitutions associated with FOP, comprising variants of the R258 residue (Hino et al., 2015; Machiya et al., 2018).

We thus performed functional characterization of p.R258W. U2OS cells were transiently transfected with both the wild-type and mutated ACVR1 cDNAs, then treated with ActA and BMP2. As shown in Figure 3, the novel variant perceives ActA as an agonist similarly to what observed for p.R206H and p.R258S, used as positive controls, whereas wild-type expressing cells are insensitive to the treatment. This effect can be observed as an increase in the activity of a BMP-responsive reporter gene, called BRE-Luc (Figure 3a);
as an up-regulation of the ID1 target gene expression (Figure 3b) and as an increase of the phosphorylation state of the SMAD1/5/9 mediators (Figure 3c and d). These data indicate that, as already reported for the other mutations associated with FOP, also the p.R258W carrying ACVR1 can transduce an aberrant intra-cellular signaling due to the altered, mutation-induced, perception of the binding with ActA. As expected, both wild-type and mutated ACVR1 constructs can elicit a response to BMP2.

4 | CONCLUSION

We described here the case of a little child with an early clinical presentation of FOP and carrier of a novel variant of the ACVR1 gene. FOP is one of the most severe genetic diseases of heterotopic ossification. Early clinical diagnosis is mandatory for the management of the patient to avoid any procedure that might be worthless and harmful. Molecular analysis with the identification of a causative mutation of the ACVR1 gene is crucial to confirm or exclude the condition, thus providing the patient and his family with the proper counseling about the nature, the course, and the management of the disease. Furthermore, it is worth to remind that FOP is an autosomal dominant condition and although most cases are sporadic and due to de novo mutations, evidence of germline mosaicism has been reported (Janoff et al., 1996) and this may be associated with a higher risk of recurrence in successive pregnancies.

So far, there is no etiologic treatment for FOP. However, many efforts are currently ongoing to develop specific therapies targeting the receptor activity and the downstream aberrant pathway or targeting cellular components and/or processes that are important in modifying the local environment promoting bone neoformation. The study of these basic mechanisms is opening the door to the development of targeted therapies (for a revision see Cappato et al., 2018; Katagiri et al., 2018) and clinical trials are presently ongoing with a RARγ agonist derivative, Palovarotene, with a humanized anti-ActA antibody and others may be active in the near future (please refer to the ClinicalTrials.gov site for a comprehensive and updated list).
Therefore, FOP patients are facing a new era with the opportunity of participating in interventional clinical studies and, hopefully, to have access to a possible therapy in the next future. The mandatory prerequisite to benefit from these opportunities is a clinical diagnosis confirmed by the molecular characterization to define the presence of a causative mutation of the ACVRI gene.

ACKNOWLEDGMENTS

We gratefully thank the patients and the FOP Italia Association of patients and their families for funding and for continuous support to our work. This work was also supported by the “Cinque per mille” and “Ricerca corrente” (Italian Ministry of Health) and developed within the framework of the DINOGMI, Department of Excellence of MIUR 2018–2022 (legge 232 del 2016).

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS

SC, functional studies, data analysis, and manuscript writing; RT, clinical diagnosis and description, patient’s clinical management, manuscript writing; JG, medical consultation, manuscript revision; FZ critical reading of the manuscript; RB, conceptualization, molecular diagnosis, manuscript writing, and revision.

ETHICAL CONSIDERATIONS

This study was performed with a written, informed consent of the proband's parents.

ORCID

Renata Bocciardi https://orcid.org/0000-0001-8415-3802

REFERENCES

Baujat, G., Choquet, R., Bouée, S., Jeanbat, V., Courouve, L., Ruel, A., Michot, C., Le Quan Sang, K.-H., Lapidus, D., Messiaen, C., Landais, P., & Cormier-Daire, V. (2017). Prevalence of fibrodysplasia ossificans progressiva (FOP) in France: An estimate based on a record linkage of two national databases. *Orphanet Journal of Rare Diseases*, 12(1), 123. https://doi.org/10.1186/s13023-017-0674-5

Bocciardi, R., Bordo, D., Di Duca, M., Di Rocco, M., & Ravazzolo, R. (2009). Mutational analysis of the ACVRI gene in Italian patients affected with fibrodysplasia ossificans progressiva: Confirmations and advancements. *European Journal of Human Genetics*, 17(3), 311–318. https://doi.org/10.1038/ejhg.2008.178

Buczko, R., Hoeman, C., Rakopoulos, P., Pajović, S., Letourneau, L., Dzamba, M., Morrison, A., Lewis, P., Bouffet, E., Bartels, U., Zuccaro, J., Agnihotri, S., Ryall, S., Barsczczyk, M., Chornenky, Y., Bourgey, M., Bourque, G., Montpetit, A., Cordero, F., … Hawkins, C. (2014). Genomic analysis of diffuse intrinsic pontine gliomas identifies three molecular subgroups and recurrent activating ACVRI mutations. *Nature Genetics*, 46(5), 451–456. https://doi.org/10.1038/ng.2936

Cappato, S., Giacopelli, F., Ravazzolo, R., & Bocciardi, R. (2018). The horizon of a therapy for rare genetic diseases: A “Druggable” future for Fibrodysplasia Ossificans Progressiva. *International Journal of Molecular Sciences*, 19(4), 989. https://doi.org/10.3390/ijms19040989

Cappato, S., Giacopelli, F., Tonachini, L., Ravazzolo, R., & Bocciardi, R. (2019). Identification of reference genes for quantitative PCR during C3H10T1/2 chondrogenic differentiation. *Molecular Biology Reports*, 46(3), 3477–3485. https://doi.org/10.1007/s11033-019-04713-x

Cappato, S., Tonachini, L., Giacopelli, F., Tirone, M., Galletti, L. J., Sormani, M., Giovenzana, A., Spinelli, A. E., Canciani, B., Brunelli, S., Ravazzolo, R., & Bocciardi, R. (2016). High-throughput screening for modulators of ACVRI transcription: Discovery of potential therapeutics for fibrodysplasia ossificans progressiva. *Disease Models & Mechanisms*, 9(6), 685–696. https://doi.org/10.1242/dmm.023929

Chang, M. T., Asthana, S., Gao, S. P., Lee, B. H., Chapman, J. S., Kandoth, C., Gao, J., Socci, N. D., Solit, D. B., Olshen, A. B., Schultz, N., & Taylor, B. S. (2016). Identifying recurrent mutations in cancer reveals widespread lineage diversity and mutational specificity. *Nature Biotechnology*, 34(2), 155–163. https://doi.org/10.1038/nbt.3391

Eresen Yazıcıoğlu, C., Karatosun, S., Kizildağ, S., Ozsoylu, D., & Kavukçu, S. (2013). ACVRI gene mutations in four Turkish patients diagnosed as fibrodysplasia ossificans progressiva. *Gene*, 515(2), 444–446. https://doi.org/10.1016/j.gene.2012.12.005

Hatsell, S. J., Idone, V., Wolken, D. M., Huang, L., Kim, H. J., Wang, L., Wen, X., Nannuru, K. C., Jimenez, J., Xie, L., Das, N., Makhoul, G., Chernomorsky, R., D’Ambrosio, D., Corpina, R. A., Schoenherr, C. J., Feeley, K., Yu, P. B., Yancopoulos, G. D., … Economides, A. N. (2015). ACVRI R206H receptor mutation causes fibrodysplasia ossificans progressiva by imparting responsiveness to activin A. *Science Translational Medicine*, 7(303), 303ra137. https://doi.org/10.1126/scitranslmed.aac4358

Hino, K., Ikeya, M., Horigome, K., Matsumoto, Y., Ebise, H., Nishio, M., Sekiguchi, K., Shibata, M., Nagata, S., Matsuda, S., & Toguchida, J. (2015). Neofunction of ACVRI in fibrodysplasia ossificans progressiva. *Proceedings of the National Academy of Sciences of the United States of America*, 112(50), 15438–15443. https://doi.org/10.1073/pnas.1510540112

Janoff, H. B., Muenke, M., Johnson, L. O., Rosenberg, A., Shore, E. M., Okerere, E., Zasloff, M., & Kaplan, F. S. (1996). Fibrodysplasia ossificans progressiva in two half-sisters: Evidence for maternal mosaicism. *American Journal of Medical Genetics*, 61(4), 320–324. https://doi.org/10.1002/(SICI)1096-8628(19960202)61:4<320::AID-AJMG4>3.0.CO;2-Y

Kailiya-Perumal, A.-K., Carney, T. J., & Ingham, P. W. (2020). Fibrodysplasia ossificans progressiva: Current concepts from bench to bedside. *Disease Models & Mechanisms*, 13(9), 6441. https://doi.org/10.1242/dmm.046441

Kaplan, F. S., Kobori, J. A., Orellana, C., Calvo, I., Rosello, M., Martinez, F., Lopez, B., Xu, M., Pignolo, R. J., Shore, E. M., & Groppe, J. C. (2015). Multi-system involvement in a severe variant of fibrodysplasia ossificans progressiva (ACVRI c.772G>A; R258G): A report of two patients. *American Journal of Medical Genetics, Part A*, 167A(10), 2265–2271. https://doi.org/10.1002/ajmg.a.37205
