Nanostring technology on Fibrous Dysplasia bone biopsies. A pilot study suggesting different histology-related molecular profiles

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A B S T R A C T
Identifying the molecular networks that underlie Fibrous Dysplasia (FD) is key to understand the pathogenesis of the disease, to refine current diagnostic approaches and to develop efficacious therapies. In this study, we used the NanoString nCounter Analysis System to investigate the gene signature of a series of nine Formalin Fixed Decalcified and Paraffin-Embedded (FFDPE) bone biopsies from seven FD patients.

We analyzed the expression level of 770 genes. Unsupervised clustering analysis demonstrated partitioning into two clusters with distinct patterns of gene expression. Differentially expressed genes included growth factors, components of the Wnt signaling system, interleukins and some of their cognate receptors, ephrin ligands, matrix metalloproteinases, neurotrophins and genes encoding components of the cAMP-dependent protein kinase. Interestingly, two tissue samples obtained from the same skeletal site of one patient one year apart failed to segregate in the same cluster. Retrospective histological review of the samples revealed different microscopic aspects in the two groups.

The results of our pilot study suggest that the genetic signature of FD is heterogeneous and varies according to the histology and, likely, to the age of the lesion. In addition, they show that the Nanostring technology is a valuable tool for molecular translational studies on archival FFDPE material in FD and other rare bone diseases.

1. Introduction

Fibrous Dysplasia of bone (FD) is a genetic skeletal disease that occurs in monostotic or polyostotic forms, the latter being often associated with extra-skeletal disorders in the McCune-Albright Syndrome (Polyostotic FD/MAS; OMIM#174800). FD results from gain-of-function mutations of the GNAS gene (GNAS complex locus; GNAS, OMIM#139320) that arise during embryo development (Riminucci et al., 2007) and usually replace the arginine 201 (R201) in the Gsα transcript with a histidine (R201H) or a cysteine (R201C) (Weinstein et al., 1991). In spite of its early occurrence, the mutation targets the osteoprogenitor cell compartment of the post-natal skeleton (Bianco and Robey, 1999; Riminucci et al., 2007) and severely compromises growing bones causing loss of mechanical strength, deformity and pain (Ippolito et al., 2014; Mancini et al., 2009). The essential pathological features of FD include fibrotic tissue, woven and hypertmineralized bone, osteoclasts and blood vessels, which may variably combine to generate site specific histological patterns (Corsi et al., 2017, 2003; Riminucci et al., 2003, 1999). However, it is known that FD lesions may also show additional microscopic aspects (e.g. cartilage, myxoid areas) (Corsi et al., 2006; Tabareau-Delalande et al., 2013) or may develop extensive secondary changes of uncertain significance (e.g. fibroma desmoplastic-like areas), especially in craniofacial bones (Corsi et al., 2020a, 2020b). In these cases, substantial deviations from the archetypal FD lesion may occur with consequent diagnostic challenges and prognostic uncertainty. Although the Gsα mutation, which is sufficient to reproduce the core pathology of FD in mice (Saggio et al., 2014), remains the essential genetic marker of the disease, a deeper knowledge of the molecular

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background is important to improve the specific diagnostic and molecular profiling of FD lesions and histological variants thereof. Previous molecular studies essentially focused on individual genes (Alman et al., 1995; Candelieri et al., 1995; Jia et al., 2011; Kashima et al., 2009; Kumta et al., 2003; Regard et al., 2011; Rimbuzzi et al., 2003; Stanton et al., 1999; Yamamoto et al., 1996), or on restricted panels of selected genes (Kiss et al., 2010) involved in skeletal growth and homeostasis. However, the complex molecular scenario downstream gain-of-function mutations of a pleiotropic gene like Gso calls for the application of high throughput methods of transcriptomic analysis and, in particular, for methods that allow a tight correlation between molecular profiling and histology. Only two microarray studies on FD are currently available. The first, performed on a pool of fresh pathological samples (Zhou et al., 2014), did not provide any correlation between the molecular data and the actual histological features of the lesions. The second did not use pathological tissue and was conducted on normal osteoprogenitor cells transduced with the GsR201C mutation (Raimondo et al., 2020).

The NanoString technology has been proposed as a novel enzyme-free methodology that allows for high throughput analysis of samples with poor extraction quality and low nucleic acid yield (Tsang et al., 2017). For this reason, it is considered as a valid approach to screen potential candidate genes on formalin-fixed paraffin-embedded (FFPE) tissue specimens, including bone (Wang et al., 2019). In this study, the expression level from 770 genes was determined by NanoString technology on a small series of formalin-fixed and paraffin-embedded (FFDPE) bone biopsies from seven FD patients.

Although limited, our cohort of samples generated two different FD molecular profiles suggesting that the genetic signature of FD is heterogeneous and varies according to the histology and, likely, to the age of the lesion. More in general, our study demonstrated that the NanoString technology is a valuable tool for molecular translational studies on archival material in FD and other rare bone diseases.

2. Methods

2.1. Patients and tissue samples

Nine archival paraffin blocks collected at the Sapienza University Hospital Policlinico Umberto I of Rome between 2008 and 2017 from seven FD/MAS patients (Pt 1-Pt 7) were used. The study was approved by the local Ethical Committee (RIF. EC: 5126). Informed consent was obtained from patients or their guardians. Five patients were males and two were females and the age range at tissue harvesting was 6–23 years. The clinical diagnosis was MAS in six patients (four males and two females) and polyostotic FD in one. For two patients two tissue samples were analyzed. The samples were obtained from femur (sample A) and tibia (sample B) for one of them (Pt 1). For the other (Pt 3), both samples (A and B) were obtained from the femur one year apart.

All tissue samples were collected during either diagnostic biopsies or corrective surgeries, immediately fixed in 4% buffered formaldehyde, decalcified with 10% EDTA and routinely processed for paraffin embedding. Mutation analysis was performed in the FD samples from six patients as described previously (Corsi et al., 2006; Rimbuzzi et al., 1999). R201H was detected in three patients (Pt 1, Pt 3 and Pt 7) and R201C in the other three (Pt 2, Pt 4 and Pt 6). A synopsis of the clinical and molecular data of the patients included in the study is reported in Table 1.

2.2. RNA extraction

Total RNA was extracted by RNeasy FFPE kit (Qiagen Germany) according to manufacturer’s instructions starting from a minimum of four 20-micron-thick sections from FFDE samples. RNA quantity and quality were assessed by NanoDrop2000 spectrophotometer (Thermo Fisher Scientific).

2.3. NanoString analysis

The samples that reached the quality standards (A260/A280 higher than 1.7 and A260/A230 higher than 1.8) were analyzed by NanoString using the nCounter PanCancer Profiling Panel (NanoString Technologies). This panel includes 770 genes from 13 canonical pathways including: MAPK, STAT, PI3K, RAS, Cell Cycle, Apoptosis, Hedgehog, Wnt, DNA Damage Control, Transcriptional Regulation, Chromatin Modification, and TGF-β. Since a specific nCounter Panel related to FD is not available to date, we selected this panel as it includes driver genes and gene pathways that, according to our experience, could provide some insights into the molecular features of FD.

Analysis of detected gene counts was performed by employing nSolver Analysis Software 3.0 (NanoString Technologies). First, samples were selected by checking imaging quality controls: percentage of fields of view (FOV) read (higher than 75%), binding density (between 0.05 and 2.25), positive control linearity (higher than 0.95), and positive control limit of detection (higher than 2). For samples that passed imaging quality controls, raw genes counts were normalized on technical controls and housekeeping genes included in the panel. Mean count of negative controls plus two standard deviations (SDs) were subtracted from each gene count to eliminate negative background. Then, normalization on positive controls was conducted by multiplying the count of each gene for a correction factor (CF) calculated, for each sample, as the ratio between the quadratic mean of positive controls counts and the mean of quadratic means in all samples.

To perform CodeSet Content normalization on housekeeping genes, ten reference genes were selected among those available in the panel based on the lowest coefficients of variation (CV = ratio between mean counts and SDs across all samples). Finally, CodeSet Content normalization was performed by multiplying genes counts for a further CF calculated on reference genes as described for positive technical controls. Expression data are available at NCBI GEO: GSE176243.

After the normalization processes, counts were log2 transformed and used for downstream analyses. To perform cluster analysis, we chose the hierarchical agglomerative clustering procedure, which has a bottom-up approach. This procedure was employed using Ward’s method and a Euclidean distance metric. Ward’s method, essentially, looks at cluster analysis as an analysis of variance problem, starting out with n clusters of size 1 and continuing until all the observations are included into one cluster. Hierarchical clustering of samples and genes was performed with the “heat map.2” R function by employing “ggplot” and “dendextend” packages. Each cell was colored to reflect quantitatively the relative expression, in NanoString counts, to assist in visualizing expression changes.

2.4. Pathway enrichment and Gene Ontology (GO) analysis

BioPlanet 2019, WikiPathways 2021 and KEGG 20121 enrichment analysis of differential gene expression in addition to Gene Ontology (GO) for biological processes, molecular function and cellular
component were performed with the publicly available tool, EnrichR (http://amp.pharm.mssm.edu/Enrichr), that provides access to various gene-set libraries. We considered pathways and Gene Ontology terms as enriched if their p-value was lower than 0.05 and sorted them by combined score ranking. The combined score is a combination of the p-value and z-score calculated by multiplying the two scores as follows: \( CS = \log(p)z \) where \( p \) is the p-value computed using Fisher’s exact test, and \( z \) is the z-score computed to assess the deviation from the expected rank. The plots show the significance \( \log(p\text{-value}) \) of each gene set from the selected library versus its odds ratio. Each point represents a single geneset. Larger blue points represent significant terms (p-value lower than 0.05) while smaller gray points represent non-significant terms.

The search tool for retrieval of interacting genes (STRING) (https://string-db.org) database, which integrates both known and predicted PPIs, has been applied to predict functional interactions of proteins (Szklarczyk et al., 2019). To search for potential interactions between overexpressed genes, the STRING tool was employed, species limited to “Homo sapiens” and an interaction score > 0.4. Cytoscape software version 3.6.1 was used to visualize the PPI network and to detect the highly connected regions of the network, Markov Clustering (MCL) algorithm was used with default parameters. In networks, nodes correspond to proteins and edges represent interactions.

### 2.5. Retrospective evaluation of the histological samples

Four-micron-thick FFDPE sections were stained with hematoxylin and eosin (HE) for transmitted light microscopy or with Picrosirius red for analysis with polarized light microscopy. TRAP staining was performed to highlight mono- and multi-nucleated osteoclasts as described previously (Corsi et al., 2020a, 2020b). The microscopic review was performed by three pathologists (MDSV, AC, MR) with reference to the FD histological patterns and sub-classification previously reported (Tabureau-Delalande et al., 2013).

Bone volume in each biopsy (Bone Volume/Tissue Volume BV/TV) was quantified on H&E stained sections according to the guidelines of the American Society of Bone and Mineral Research (Dempster et al., 2013). For the comparison of BV/TV between the two groups, the two-sided unpaired Students’ t-test was employed. Statistical significance was established at p-value lower than 0.05.

### 3. Results and discussion

#### 3.1. NanoString analysis revealed different FD molecular clusters

We isolated total RNA from 9 archival FFDPE FD bone lesions [two samples (A and B) from Pt1 and Pt3 and one sample from Pt2, Pt4, Pt5, Pt6 and Pt7]. The RNA yield was adequate in all cases and an RNA input between 150 ng and 250 ng was used for subsequent NanoString analysis. Two samples (Pt1 A and Pt6) were excluded due to degradation of RNA as detected by preliminary analysis of raw data performed by nSolver™ software. In fact, they presented Quality Check flag for “content normalization” indicating an effective reduction in quality (fragmentation) of the input RNA.

Unsupervised clustering analysis demonstrated that the remaining 7 FD samples could be clearly partitioned into two groups (defined FD cluster 1 and FD cluster 2) with distinct patterns of gene expression (Fig. 1). Interestingly, the two tissue samples obtained from the same skeletal site of Pt3 one year apart during the progression of the disease.
did not segregate in the same group.

To confirm the reliability of the Nanostring approach, we randomly selected 5 genes that resulted highly expressed and 5 genes that showed low levels of expression and tested their expression levels by quantitative PCR (qPCR) on two frozen FD bone samples. As reported in Fig. S1, the results of the qPCR analysis were consistent with the data obtained by Nanostring. The sequence of the primers used for qPCR is reported in Table S2.

3.2. Differentially expressed genes and pathway enrichment

FD cluster 1, which included samples from Pt2, Pt3 (sample A) and Pt4, was characterized by the upregulation of 189 genes (Table S1). These included: growth factors; components of the Wnt signaling system; interleukins and some of their cognate receptors; ephrin ligands; matrix metalloproteinases and neurotrophins. Interestingly we also observed upregulation of genes encoding components of the cAMP-dependent protein kinase as the catalytic subunit gamma (PRKACG) and the type I-beta regulatory subunit (PRKAR1B) (Table S1).

Fig. 2. Pathway enrichment and Gene Ontology (GO) analysis pathway enrichment analysis. Bioplanet 2019 (A, B), WikiPathways 2021 (C, D) and KEGG 2021 (E, F) pathway enrichment analyses of the 188 differentially expressed genes. Pathways in A, C, E are sorted by combined score ranking. Plots in B, D, F show the significance [−log10(p-value)] of each gene set from the selected library versus its odds ratio. Each point represents a single geneset. Blue points represent significant terms (p-value lower than 0.05) while gray points represent non-significant terms.
transmitted light microscopy. B: polarized light microscopy. trabeculae (D and E) and a very low number of orthotopic osteoclasts. A, D and E: hematoxylin and eosin. B: Picrosirius red. C and F: TRAP. A, C, D, E and F: red staining in C). In contrast, the lesions included in molecular cluster 2 were characterized by a larger amount of fibro-myxomatous tissue with sparse bone lesional cells is known to be associated with abnormal osteoblast differentiation and formation of defective bone. WNT signaling pathways, known to regulate bone mass and development (Liu et al., 2008), are enriched in cluster A.4.

Moreover, functional enrichment analyses, including BioPlanet 2019, WikiPathways 2021 and KEGG 2021 databases at Enrichr (http://amp.pharm.mssm.edu/Enrichr), demonstrated that Cytokines and inflammatory response, PI3K-AKT and MAPK signaling pathways, and FGFR ligand binding and activation were the top enriched pathways (Fig. 2).

### 3.3. Correlation between molecular clusters and FD histology

Retrospective evaluation was performed with reference to the histological subtyping proposed by Flore Tabareau-Delalande et al. who distinguished conventional and unconventional histological patterns of FD (Tabareau-Delalande et al., 2013). All lesions that segregated in the FD cluster 1 [Pt2, Pt3 (sample A) and Pt4] met the criteria of conventional FD. In these samples, histological sections were occupied by a highly vascularized fibrotic tissue and haphazardly distributed pathological bone trabeculae with a woven structure and well evident Sharpey’s fibers. In all cases, osteoclasts were promptly recognized in close contact with the FD bone. Increase number and thickness of FD bone trabeculae and heterotropic (i.e., within the fibrotic tissue) osteoclasts were observed in the samples from Pt4 and Pt2, respectively.

In contrast, the samples that segregated in the FD cluster 2 [Pt1 (sample B), Pt 3 (sample B), Pt5 and Pt7] shared an unconventional fibro-involutive pattern. A reduction in the amount of bone, even though not statistically significant was observed compared to FD molecular cluster 1 (BV/TV, mean ± SD, molecular cluster 1 vs molecular cluster 2, 28.87 ± 3.07 vs 23.52 ± 11.83). Focal areas of conventional FD with rare, if any, osteoclasts in contact with FD bone trabeculae were detected in samples from Pt 3 (sample B), Pt5 and Pt7.

Representative histological images of the FD lesions segregating in the FD molecular cluster 1 [from Pt 3 (sample A)] and in FD molecular cluster 2 [from Pt 1 (sample B)] are illustrated in Fig.S3. Additional histological features of both clusters are reported in Fig. S2.

Interestingly, the two samples obtained from Pt3 one year apart did

![Histological Images](image-url)

**Fig. 3.** Representative histological images from FD lesions segregating in molecular cluster 1 [A–C, Pt 3 (sample A)] and in molecular cluster 2 [D–F, Pt 1 (sample B)]. The lesions included in molecular cluster 1 consisted of a highly vascularized fibrotic tissue (ft) and haphazardly distributed bone trabeculae (bt) (A) and showed well evident Sharpey’s fibers (arrows in B) and a large amount of orthotopic (i.e., along the bone surfaces) and heterotopic (i.e., within the fibrotic tissue) osteoclasts (red staining in C). In contrast, the lesions included in molecular cluster 2 were characterized by a larger amount of fibro-myxomatous tissue with sparse bone trabeculae (D and E) and a very low number of orthotopic osteoclasts. A, D and E: hematoxylin and eosin. B: Picrosirius red. C and F: TRAP. A, C, D, E and F: transmitted light microscopy. B: polarized light microscopy.
not segregate in the same group. The changes in the overall histological and molecular features of these two samples could result, at least in part, from age-dependent changes in the biology of the FD tissue. For example, we have previously shown that a reduction in the number of mutated osteoprogenitor cells occurs within FD lesions over time eventually leading to a quiescent state (Kuznetsov et al., 2008).

4. Conclusions

This is a very preliminary study with multiple limitations, as the reduced number of pathological samples and the use a NanoString nCounter Kit not specific to investigate bone diseases. Nonetheless, our results reveal that the genetic signature of FD, as its histology, is heterogeneous and likely age-dependent and provide some relevant data for future molecular translational research. Furthermore, they indicate that the NanoString technology is a valuable tool for molecular studies on archival histological FFPE material from rare bone diseases.

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Declarations of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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