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CFTR/CEREBRAL STRESS

A1

Myriocin potential as a phenotype-modifying therapeutic in Cystic Fibrosis

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Background

Cystic Fibrosis (CF) is caused by a variety of mutations of the CFTR ion channel, with deletion of phenylalanine 508 (F508del) representing approximately 70% of patients. This mutation induces a proteinopathy, characterized by aggregates of mutated and unfolded proteins, hyper-inflammation, impaired trafficking and altered metabolism at cellular level. CFTR dysfunction has primarily a devastating effect on lungs, causing chronic inflammation and recurrent unresolved infections. CF patients develop severe comorbidities: male infertility, biliary cirrhosis, osteopenia/osteoporosis, renal failure, diabetes (50% of adult patients), malabsorption and increased synthesis of cholesterol and its accumulation in liver but also in other tissues, dyslipidemia with increased plasma triglycerides, pancreas fibrolipomatosis, hepatic lipogenesis and steatosis. Lung chronic inflammation and infection have been extensively associated to the lipotoxin ceramide, core component of membrane lipids. such as sphingomyelins and glycosphingolipids. We previously demonstrated that the sphingolipid synthesis inhibitor Myriocin is able to reduce inflammation and to ameliorate defense response against bacterial and fungal infection.

Materials and methods

43 CF patients with selected genotypes were enrolled. Monocytes from peripheral blood samples were extracted and also clinical data were collected.

Results

We here demonstrate the mode of action of this molecule. First, we show in IB3 cell line, that Myriocin is an effective inducer of autophagy which is defective in CF proteinopathy and can sustain pathogen clearance (xenophagy). Second, we demonstrated that Myriocin activates key transcriptional factors, TFEB, FOXO1a and PPARgamma involved in autophagy induction, mitochondria genesis and activity, energy production and lipid mobilization and consume. We proved that Myriocin significantly increases the transcription of downstream genes regulating fatty acids entry in mitochondria (CTP1a and 1b; FATP) and their oxidation (ACAD L). Next, we showed that Myriocin dramatically reduces pathological accumulation of lipid unorganized deposits in IB3 cells. By Mass spectrometry we observed that inhibition of sphingolipid synthesis causes a reduced content of non sphingoid-base containing lipids, such as glycerol lipids and cholesterol. We then proved that Myriocin is able to change the transcriptional profile of IB3 treated cells, by gene sequencing analysis, enhancing the transcription of genes involved in lipid transport and consume and energy metabolism that resulted partially downregulated in CF. Finally, Myriocin treatment of peripheral blood monocytes from CF patients, infected with A. fumigatus conidia, significantly increases their pathogen killing ability.

Conclusions

Myriocin is a potent modifier of pathological gene expression profile in CF and a potential therapy for CF related infection.

A2

miR-125b/NRF2/CFTR circuitry: a pilot study on oxidative stress in cystic fibrosis

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Background

In physiopathology of Cystic Fibrosis (CF), oxidative stress implications were recognized and widely accepted. The CFTR defects in ionic transport disrupt the intracellular redox balance causing CF classical pathologic hallmarks. Therefore, oxidative stress together with detoxification genes and miRNA aberrant expression could be associated with clinical outcome.

Materials and methods

Patients (n=8) with CF diagnosis with the same mutation (F508del), but various clinical status, compared to healthy individuals. After nasal brushing, we extracted total RNA of patients for consecutive expression analysis of CFTR, NRF2 and its targets as well as miR-125b, by quantitative real time PCR (qPCR) using TaqMan chemistry. Written informed consent was obtained by each participant to the study.

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Results
In this pilot study, we analyzed the existence of a correlation among CFTR, miR-125b, NRF2 and oxidative stress genes in a representative number of CF patients. In CF patients with chronic P. aeruginosa (PA) lung infection, the mRNA levels of the NRF2 gene were significantly down-expressed and showed a direct correlation with CFTR gene expression. Interestingly, the NRF2 downmodulation was accompanied by an induced expression of miR-125b. On the contrary, PA negative CF patients showed NRF2 expression levels more similar to those of healthy individuals, with a reduction of miR-125b, even if CFTR levels remained downmodulated. These data are in line with the expression of the oxidative stress target genes NQO1 and GST-T1. Moreover, we found that PA positive patients with a FEV1>60% showed the expression of HMO1, another NRF2 target gene, higher than those with a lower FEV1.

Conclusions
The evidence of a CFTR, NRF2 and miR-125b impaired network as oxidative stress response in CF patients, prompt us to hypothesize that these molecular mechanisms could explain the wide CF phenotype variability as an additional control level over the CFTR gene mutations. Furthermore, this study may allow the discovery of potential therapeutic targets in order to improve CF patient’s quality of life by screening the expression of these oxidative stress factors as prognostic markers.

MODEL SYSTEMS

A3
CFTR-Dependent Bicarbonate Transport in Human Rectal Biopsies carrying CFTR variants
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A3

Background
Intestinal Current Measurements (ICM) have been utilized for several decades for measuring ex vivo CFTR function as support for difficult diagnosis and for detection of effects of CFTR modulators in preclinical studies, as well as for monitoring therapies of patients affected by cystic fibrosis (CF). Transport of both chloride and bicarbonate is mediated by CFTR, a transmembrane channel permeable to anions. Defective bicarbonate transport plays a role in CF and CFTR related disorders leading to pancreas and lung damage. Selective defect of bicarbonate transport was previously reported for a group of CFTR variants in patients with pancreatitis and no lung disease (LaRusch et al., PLOS Genetics 2014; Park et al., Gastroenterology 2010). We set up ICM in appropriate conditions for detecting selective CFTR mediated/regulated bicarbonate transport defect.

Materials and Methods
We performed ICM in rectal biopsies of 8 non CF subjects, 8 CF patients and 8 patients with unknown diagnosis referring to the Cystic Fibrosis Center of Verona for CFTR function measurements. ICM were performed according to both the procedure of European Cystic Fibrosis Society ICM Standard Operating Procedure and variations using CI free or HCO3 free medium and carbonic anhydrase/transport inhibitors in non CF controls vs. wide spectrum of CFTR mutations including CFTR bicarbonate defective mutants.

Results
Normal tracings for both anions were observed in non CF controls while abnormal tracings in terms of Cl and bicarbonate transport were obtained in all CF patients except one with Q143H/R553X/R806G CFTR mutation with pancreatitis with pancreas divisum and lung infection (Staphylococcus aureus methicillin resistant). In subjects with inconclusive diagnosis in the presence of disseminated bronchiectasis we found normal tracings for both bicarbonate and Cl except in two patients with selective defect of bicarbonate transport in the presence of I807M+/-, R668C+/-, CFTR genotypes and Cl sweat values of 18 and 23 mmol/L and 28 and 68 mmol/L, respectively (sweat test by Gibson and Cooke performed twice). None had clinical history of pancreatitis in addition to lung disease, evaluation is in progress.

Conclusions
Selective defect of bicarbonate transport might be responsible for clinical phenotypes milder than classical CF, including not only pancreas, but also lung disease. It can be detected by bioassays such as ICM that is very relevant for understanding functional impact of CFTR variants and their response to CFTR modulators. Informed consent was obtained from all subjects.

Study supported by: Italian Cystic Fibrosis Research Foundation grant #7/2016, Lega Italiana FC-Associazione Veneta Onlus.

A4
Anti-estrogen and direct effects of GB-1877 on calcium-activated chloride channels
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Background
Although the median age of survival is increased in Cystic Fibrosis (CF) patients, females continue to have a lower median age of survival compared with males. Several lines of evidence indicate that estrogens play a relevant role in the pathophysiology of CF. In vitro studies have shown that 17β-estradiol inhibits airway surface liquid production, and in vivo studies in CF animals have demonstrated that 17β-estradiol enhances inflammation, increases mucus production and favors the conversion from the non-mucoid to the mucoid form of P. aeruginosa. The aim of this study was to investigate the effects of 17β-estradiol and GB-1877 on calcium-dependent chloride channel (CaCC) currents in human bronchial epithelial cells carrying the F508del-CFTR mutation both in homozygosis and in heterozygosis.

Materials and Methods
Perforated patch clamp experiments were performed on single cells using two immortalized cell lines, CFBE (F508del/F508del) and IB3-1 (F508del/IV1282X). Gramicidin (10 or 20 μM) was added to the electrode solution to reach the whole cell configuration. Electrical stimulation protocol consisted in voltage steps from -80 to +80 mV with 20 mV interval and 800 msec duration.

Results
The presence of 17β-estradiol significantly reduced the CaCC currents, both in basal conditions and in the presence of ATP (100 μM). The addition of GB-1877 (10 μM) completely restored the currents abolished by 17β-estradiol, in basal conditions and after stimulation with ATP in both CFBE and IB3-1 cells. GB-1877 had a strong, direct action on membrane current density, which significantly increased more than 4-fold in both cases. The membrane current stimulation produced by GB-1877 was further enhanced by the addition of ATP. CFBE cells incubated for 24 hours with 3 μM VX-809 (a CFTR corrector) and then acutely stimulated with VX-770 (a CFTR potentiator) in the presence of forskolin, showed an increase of chloride currents.
which were abolished by Inh-172. The chloride current density induced by GB-1877+ATP was, on average, greater than that obtained with VX-809+VX-770+ forskolin. The currents elicited by GB-1877+ATP were abolished by the addition of NPPB, a CaCC inhibitor, indicating that the membrane currents recorded in CFBE and IB3-1 cells were mainly carried by chloride through the CaCC. The combined administration of GB-1877+ATP and VXs/FSK had an additional effect on chloride currents.

Conclusions

Our results show that GB-1877 restores CaCC currents inhibited by 17β-estradiol and directly activates the CaCC-dependent chloride currents potentiated by ATP, an effect which is mutation independent. The combined effect of GB-1877 with currently used treatments for CF (e.g., CFTR correctors and potentiators, ENaC inhibitors) could be of benefit to patients with CF.

A5

Quantification of CRISPR/Cas9 CFTR gene editing events using droplet digital PCR

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A5

Background

In the last few years, the field of therapeutic gene editing is rapidly developing with the emergence of CRISPR/Cas9 system. This developing technology represents a new opportunity for the treatment of many genetic diseases, including Cystic Fibrosis. However, despite the great potential of this technique, the low frequency of genome editing events, along with the off-target events generation, remains a limitation in the clinical application of this powerful method. In an attempt to develop methods to improve the efficiency of genome editing events, we set up a protocol for quantifying these editing events using the droplet digital PCR (ddPCR) technology.

Materials and methods

For this purpose, the ddPCR has already been performed using labeled probe (i.e. fam, hex or vic) technology. Our method, based on EvaGreen technology, uses two primer pairs: one primer pair amplifies a reference sequence distant from the Cas9 cleavage site (reference assay); and a second primer pair include the Cas9 cleavage site (Editing assay). When a reporter expression cassette, included in the HDR construct, is inserted in the Cas9 cleavage site, the amplification of the editing assay is blocked as the ampiclon length far exceeds 1000 bp, resulting in a reduction of positive droplets from the two assays (reference and editing one) is possible to calculate the percentage of edited allele. The quantification of the genome editing events, using the ddPCR assay, was performed on gDNA of 293T cells transfected with two constructs: 1) one home-made HDR construct (bearing a reporter cassette including the Cas9 exons 11), 2) one construct expressing the Cas9 enzyme and the gRNA targeting the genomic sequence of F508del mutation.

Results

The combination of the reference and editing assay resulted in a fine quantification of gene editing events. Indeed, we were able to accurately quantify the edited alleles in 293T cells transiently transfected and after puromycin selection.

Conclusion

Finally, we demonstrated that the Droplet Digital PCR assay, based on EvaGreen technology, represent an easy and cost effective assay to evaluate the genome editing efficiency, showing in our case, a genome editing efficiency up to 90% of alleles in puromycin selected cells.

We thank Ministero della Salute (Rome, Italy) L.548/93 for the regional research funding quote 2012-2015.

A6

The role of functional studies in the diagnosis and treatment of cystic fibrosis: comparing the case of the G970D and G970R mutation

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A6

Background

More than 2000 CFTR mutations have been identified so far, but only few of them are clearly defined as CF-causing based on functional studies. We present a case of a rare mutation, the G970D, that has been shown using transfected cDNA in HEK293 cells to be sensitive to ivacaftor. However, a similar missense mutant, G970R in the same codon, was found to be sensitive to potentiators in vitro but not in vivo due to splicing alteration. Thus, we used several basic research methodologies to evaluate if this patient was eligible for treatment with ivacaftor.

Materials and Methods

1) Nasal epithelial cells (HNEC) were collected from patients to evaluate the effect of mutations on splicing by RT-PCR assay, 2) HNEC were expanded and polarized for evaluation of CFTR function by using chamber system 3) the use of a minigene system was used to confirm in vitro the splicing pattern.

Results

Firstly, we used in silico tool to predict the fisio-pathological effect of mutations, confirming that the G970R completely abolishes the canonical 5’splice donor site of exon 17 resulting in a likely retention of intron 17. On the contrary, the G970D predicted not to affect splicing. This prediction was confirmed by RT-PCR analysis of mRNA extracted from HNEC cells and from in vitro minigene assay. Finally, the functional behavior of CFTR, from HNEC bearing the G970D, was evaluated by short-circuit recordings. The cells responded to cAMP agonist with an increase in trans-epithelial current and this current was nearly doubled by stimulation with ivacaftor. Moreover, in cells that were also incubated with VX-809 (1 μM) for 24 hours, CFTR function was significantly enhanced, with a proportional increase of both cAMP- and potentiator-dependent responses.

Conclusion

Our results show that the G970R actually disrupts the RNA splicing thus leading to a severely altered CFTR protein. This event explains the lack of success of treatment with potentiator. In contrast, the G970D does not alter RNA splicing. The resulting G970D mutation affects the gating and trafficking of the CFTR channel that can be targeted with VX-770 and VX-809. These results represent the evidence needed to justify the treatment of the patient with these drugs. Finally, our study is an interesting example of the precision medicine approach by emphasizing the role of appropriate in vitro studies, in this case focused on RNA analysis, to fully characterize the effects of rare CFTR mutations.

We thank Telethon Foundation (TMLGCBX16XT) and Ministero della Salute (Rome, Italy) L.548/93 for the regional research funding quote 2008-2015.
GENETICS

A7

Description of Clinical Cases of Disease CFTR Correlated/Atypical Cystic Fibrosis. Genotype/Phenotype Correlation in Patients with Polymorphism TG12-13 ST

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A7

Background

Cystic Fibrosis (CF) is the most common genetically determined, life-limiting disorder in populations of European ancestry. The genetic basis of CF is well established to be mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene.CF is a multisystem disease affecting organs where CFTR is expressed (airways, gastrointestinal and reproductive tract).

According to the literature data, variant R31C does not cause CF when combined with another CF-causing variant. Most individuals with this variant (combined with another CF-causing variant) will be healthy. A small number of individuals may develop mild symptoms or be diagnosed with a CFTR-related disorder (CFTR-RD), but symptoms are not expected to be severe enough to meet the definition of CF. There are two scenarios in which making a diagnosis after a positive NBS is less easy:

- borderline sweat test (30-60 mmol/L) without gene mutation;
- normal sweat test with two mutations, at least one of which is of uncertain significance.

There has been a recent Delphi consensus process upon which our management is based and the terminology agreed of CF Screen-positive, Inconclusive Diagnosis (CFSPID).

Case report

G.S., male, 5 years. No familiarity with genetic diseases. Full term born, meconium issued in the first 24 hours of life. Positive screening for CF. 1 level of genetic test was negative for the 32 mutations examined, but presence of polymorphic TG12 ST site, considered "mild mutation". He performed several sweat tests in stable conditions with a doubtful or inconclusive result. Sequencing of the CFTR gene showed the presence in exon 2 of the R31C mutation in heterozygosis. The patient had a good staturo-weight growth, normal pancreatic function, mild respiratory symptoms; radio-diagnostics tests show minimal lung lesions. Culture of the sputum shows first Staphylococcus aureus, then Pseudomonas aeruginosa, treated with antibiotic therapy until eradication, and respiratory therapy.

Conclusion

The R31C mutation, according to the literature data, is defined as "Non CF causing", therefore without giving symptoms of disease if not associated with "serious" mutations. Our case is a variant R31C associated with a "mild" mutation (TG12-5T) that presents a clinical suggestive of CF. We can exclude therefore that this mutation, even associated with non-serious mutations, cannot develop a CFTR-RD picture with mild symptoms or, in males, a possible sterility secondary to agenesis of deferent ducts? What happens in children diagnosed as CFSPID? Many will remain asymptomatic, 10-20% instead will develop symptoms suggestive of CF (for example PA infection), but the trend is still under study.

Parents gave consent to patient data publication.

A8

Genotype and phenotype in a patient with R31C and TG12-5T

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A8

Background

We present two pediatric clinical cases with diagnosis of Cystic Fibrosis (CF) based on the presence of one mutated allele, F508del, and clinical symptoms. The first case, a female born in 2001 (patient 1), was diagnosed for symptoms at the age of 11 months. The sweat concentration of chloride showed clearly pathological values. However, sweat tests sometimes negative/border line, the average of our case studies showed variable chloride concentration values, suggesting the presence of a CFTR polymorphism associated with classical mutation responsible for CF. I level of genetic test was negative for the 32 mutations examined, but presence of polymorphic TG12 ST site, considered "mild mutation". He performed several sweat tests in stable conditions with a doubtful or inconclusive result. Sequencing of the CFTR gene showed the presence in exon 2 of the R31C mutation in heterozygosis. The patient had a good staturo-weight growth, normal pancreatic function, mild respiratory symptoms; radio-diagnostics tests show minimal lung lesions. Culture of the sputum shows first Staphylococcus aureus, then Pseudomonas aeruginosa, treated with antibiotic therapy until eradication, and respiratory therapy.

Conclusion

The R31C mutation, according to the literature data, is defined as "Non CF causing", therefore without giving symptoms of disease if not associated with "serious" mutations. Our case is a variant R31C associated with a "mild" mutation (TG12-5T) that presents a clinical suggestive of CF. We can exclude therefore that this mutation, even associated with non-serious mutations, cannot develop a CFTR-RD picture with mild symptoms or, in males, a possible sterility secondary to agenesis of deferent ducts? What happens in children diagnosed as CFSPID? Many will remain asymptomatic, 10-20% instead will develop symptoms suggestive of CF (for example PA infection), but the trend is still under study.

Parents gave consent to patient data publication.
tests were positive (80 and 92 mmol/L) and she presented pancreatic insufficiency, pseudo-Bartter syndrome (loss of salts) at 1 years old and portal hypertension with esophageal varices. Her last forced expiratory volume in 1 second (FEV1) resulted to be 108% of predicted. She presents chronic colonization by S. aureus and intermittent by P. aeruginosa. The second case, a male born in 2012 (patient 2), was diagnosed by newborn screening with positive value of immunoreactive trypsinogen (IRT, 69 ng/mL), the detection of F508del mutation and positive sweat tests (95 and 102 mmol/L). He presented pancreatic insufficiency and two episodes of distal intestinal obstruction syndrome. His last FEV1 resulted 96% of predicted. Also the patient 2 presents chronic colonization by S. aureus and intermittent by P. aeruginosa. To complete the genotypes of both patients, an extensive genetic and functional analysis of Cystic Fibrosis Transmembrane conductance Regulator (CFTR) was performed.

Materials and methods
DNA samples were extracted from peripheral venous blood and analyzed by DNA sequencing with ABI3130xl Genetic Analyzer. RNA samples, obtained from nasal brushing of patients, were extracted, reverse transcribed and amplified by RT-PCR. Results were analyzed by a semiquantitative densitometric assay. The DNA and RNA amplicons were extracted from agarose and sequenced.

Results
In both patients, the genetic characterization confirmed the presence of F508del mutation. The analysis of the exon 9 and surrounding intronic regions of CFTR showed an extra ampiclon larger (of about 300 nucleotides) than the wild-type. The DNA sequence revealed, in both patients, the presence of an insertion of part of intron 9 in intron 8 of the CFTR gene, within the (TG)m repeat, with a poly-T stretch. The molecular lesion resulted to be the same in both patients, with only a small difference in the number of T in the poly-T stretch. The functional characterization at RNA level revealed a near complete skipping of the mutated allele of both patients. Consequently, the mutated allele is expected not to contribute to the formation of functional CFTR protein.

Conclusions
This alteration has not been previously described in the literature. Its molecular and functional features are compatible with the definition of new CF-causing mutation of the CFTR gene. This allowed the completion of the genetic characterization of both patients. The fact that the only difference in the two mutated alleles is the small change in the number of T within the poly-T stretch allows speculating about the molecular mechanism of divergence from a common ancestral allele. We declare to have the informed consent statement for the process-number of T within the poly-T stretch. The fact that the molecular and functional features are compatible with the definition of both patients. Consequently, the mutated allele is expected not to contribute to the formation of functional CFTR protein.

Materials and methods
We produced in E.coli an engineered version of PON2, with activities matching those described for the native protein, used to generate mutants to understand its structural and biochemical details. We set up qRT-PCRs to highlight the presence of the most abundant mRNA isoforms and to genotype for the two most common PON2 SNP coding variants p.Ala148Gly and p.Ser311Cys. We investigated the regulation of expression of PON2 mRNA, by checking an hypothesis of control via an "mRNA operon", through the silencing of target genes.

Results
We focused to identify the post translational modification and we showed a 3OxoC12-dependent ubiquitination (UBQ) (LYS 168) of PON2 nearby a polymorphic site, able to influence the lactonase activity. Recently a second UBQ site, nearby a second polymorphism in position 311, has been identified by us; then we have produced mutants of the two PON2 polymorphic sites (148A/G and 311 S/C) to study the activities and relationships with PTMs. Studying the mechanism of control of PON2 regulation, we identified a RNA binding protein and a E3 ubiquitin ligase that, when silenced, is able to increase the expression of the PON2 gene. Moreover we genotyped for the SNP 311 several cell lines identifying normal, heterozygotes and also mutant homozygotes.

Conclusions
Considering the impact of this gene on the defence against gram-negative, its involvement in the inflammation, the indication that the polymorphisms could dictate the severity of disease -as already demonstrated for other pathology- it could be useful to analyze CF patients to understand if the gene can have a role of modifier in relation to the presence of SNPs.

NEW THERAPIES AND OUTCOME MEASURES

A11
Cystic fibrosis and new therapeutic strategies: use of Ivcavator in a pediatric patient
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A11

Background
Cystic fibrosis (CF) is the most common genetically determined, life-limiting disorder in populations of European. The genetic origin of CF is the mutations in the CF transmembrane conductance regulator (CFTR). CFTR mutations may be classified into 6 different categories based on the mechanisms that are affected. Class I result from mutations leading to absent CFTR production. Class II, including Phe508del, are caused by defective CFTR processing. Class III mutations result in expression of CFTR, the channel gating is defective and results in impaired chloride transport function. Conductance defects are seen in class IV mutation. Class IV result in a milder phenotype. Finally, class VI mutations are characterized by a functional but unstable CFTR [1]. Ivacaftor is a potentiator that augmented chloride transport and increased airway surface liquid height and cilia beat frequency in airway epithelial cells expressing a CFTR gating mutation (II Class) [2].

Case report
C.N. female. 5 years old. Born at the end of gestational age, birth weight 3600 g. Diagnosis by screening and familiarity with FC. Sweat test: 98 mEq/L. CF. Genetics: F508del/G1244E. Intermittent colonization by Pseudomonas aeruginosa and Staphylococcus aureus. Growth curve under 25th centile from the first months of life. She has only 6 months of follow-up. She assumes Ivacaftor at a dose of

A10
Understanding the network of the regulation of PON2 expression
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A10

Background
3OxoC12 acylhomoserine lactone (3OxoC12) is the master regulator of Quorum Sensing (QS) regulating the expression of several bacterial virulence. Paraoxonase2 (PON2) has lactonase and antioxidant activities that are inhibited by 3OxoC12. The functional characterization at RNA level revealed a near complete skipping of the mutated allele of both patients. Consequently, the mutated allele is expected not to contribute to the formation of functional CFTR protein.

Conclusions
This alteration has not been previously described in the literature. Its molecular and functional features are compatible with the definition of new CF-causing mutation of the CFTR gene. This allowed the completion of the genetic characterization of both patients. The fact that the only difference in the two mutated alleles is the small change in the number of T within the poly-T stretch allows speculating about the molecular mechanism of divergence from a common ancestral allele. We declare to have the informed consent statement for the process-number of T within the poly-T stretch. The fact that the molecular and functional features are compatible with the definition of both patients. Consequently, the mutated allele is expected not to contribute to the formation of functional CFTR protein.
75 mg every 12 h with good compliance. From the beginning of the therapy program, an improvement of the auxometric parameters and of the score of the CFQ-R respiratory domain was observed; she did not present respiratory exacerbations and showed a good tolerance to the drug, in fact no adverse events were reported. There was not enough compliance to perform the spirometry, necessary for the assessment of respiratory function.

**Conclusion**
Kalydeco is an example of innovative therapeutic strategy for carriers of a CFTR channel gating mutation. The further development of such approaches offers great promise for future therapeutic strategies in CF [3].

Parents gave consent to patient data publication.

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**Table 1 (abstract A12). Results**

| Patient | Duration of IVA (weeks) | Mean data during IVA | Mean data during follow-up | Mean absolute change |
|---------|-------------------------|----------------------|---------------------------|---------------------|
| 1       | 13                      | 14.9                 | 13.9                      | +1.0                |
| 2       | 13                      | 14.4                 | 14.5                      | +0.1                |
| 3       | 13                      | 14.0                 | 14.0                      | +0.0                |

**A13**

**Long-term effectiveness of Ivacaftor in patients with Cystic Fibrosis carrying CFTR mutations with residual function and severe lung disease**

**Background**
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**Italian Journal of Pediatrics 2019, 45(Suppl 1):A13**

**Background**
Ivacaftor is a CFTR potentiator approved for class-Ill and R117H CFTR mutations and, in the US, for other 28 mutations with “residual function” (RF) of CFTR. RF mutations are usually associated with lung disease that can be delayed in onset and slower in progression, but it may become severe in the adult age.

**Materials and methods**
We describe the effectiveness and safety of Ivacaftor in subjects with CF and severe pulmonary function, at least one RF mutation. 10 subjects (Male = 1) were included (Median age 48.3 yrs, range 21.7-58.1 yrs). Genotypes were F508del/3849+10K(C-T) (n=3), F508del/D579G (n=2), D110H/D1152H (n=1), N1303K/D1152H (n=1), ES585X/3272-26A-G (n=1), Dele0x22-23/R352Q (n=1), N1303K/R117C. Ivacaftor 150 mg bid was given as compassionate use after approval by the
local Ethics Committees. Lung function, use of antibiotics, nutrition, and microbiology were evaluated before starting Ivacaftor and meanly every 3 months in the follow up.

Results
Treatment with Ivacaftor resulted in improvement of lung function. The mean absolute change (MAC) in ppFVC and ppFEV1 from the baseline value was 7.9% (95% confidence interval [CI], 4.6 to 11.2) and, respectively, 6.5% (95% CI 4.1 to 8.9) after 4 weeks (w). At 24w, the MAC of predicted FVC and FEV1 was 16.8% (95% CI 9.4 to 24.2) and, respectively, 12.7% (95% CI 3.4 to 21.9) and at 52w was 12.6% (95% CI 4.6 to 20.5) for FVC and 9.6% (95% CI 3.8 to 15.4) for FEV1. Total days of antibiotic therapy dramatically decreased from 480 days (IV 152) in the 6 months before the treatment to 138 days (IV 15) in the 12 months of follow up. No hospitalization was recorded in the follow up. Mean (SD) BMI increased from 23.1 kg/m² (5.8) at baseline to 23.5 (4.3) at 24w and to 25.5 (4.3) at 52w. Sputum microbiology was unchanged. No safety concerns were registered.

Conclusions
This case series expand our knowledge about potential benefits of Ivacaftor for CFTR mutations with CF, showing significant and sustained improvement of lung function and BMI and a considerable decrease of pulmonary exacerbations.

A14
CFTR Modulators effect on physical performance: preliminary data from CF Tuscany casuistry
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A14

Background
CFTR modulators and potentiators opened a new scenario in the treatment of specific genetic defects among CF patients. These new therapeutic options are still undergoing a regulatory and distribution process by Italian healthcare system to become part of drugs portfolio for CF patients. CF Centers throughout Italy were able to prescribe these drugs firstly by a special compassionate use program (CUP) tailored for severely compromised patients and secondly by a regulated program of prescription, administration and monitoring of responsiveness, adverse effects and adherence.

Materials and methods
Between July 2014 and September 2018 at the Regional Reference Center for the CF of Florence, a total of 45 patients, 26 females, mean aged 30.4 (6–54) years with CF, carriers of a CFTR channel gating mutation, were enrolled and assigned to receive CFTR modulator therapy with Ivacaftor (IV) or a combination of Ivacaftor and Lumacaftor (IV-Lu). For IV-Lu group, among other medical parameters, spirometry and six-minute walking test (6MWT) were performed at baseline and after 12-month as functional markers of responsiveness. Data were expressed as mean (range).

Results
In our centre to date, 12 patients started IV, 75% of which commenced it with CUP. IV-Lu therapy involved 33 patients, 36% of patients started with CUP. At baseline mean FEV1 % of predicted of IV group was 61.5 (23-105) and for IV-Lu group 67.5 (16-161). In both groups the most used airway clearance technique was positive expiratory pressure mask and only 4 patients used periodic CPAP, both techniques were performed twice a day. Mean period of therapy was 24 months for IV group and 13 months for IV-Lu group. By now 15 IV-Lu patients (45%) completed 12-month of treatment. Among them mean absolute difference of distance walked at 6MWT was -15.2 (+82, -191) meters. 7 patients showed an improvement in 6MWT distance reaching the minimal important difference (45 meters) only in two cases. Also percentage of predicted values of 6MWT distance showed no significant improvement.

Conclusions
The high heterogeneity of our casuistry related to respiratory disease and functional capacity underlines the need for a personalized approach to assess the responsiveness to these new drugs. To date data were not available for the entire sample, nevertheless we speculate that a close monitoring of adherence to respiratory physiotherapy should be mandatory and 6MWT distance could be an inaccurate outcome for IV-Lu responsiveness without the contribution of other more reliable outcomes of maximal exercise performance.

A15
Effects of Lumacaftor/Ivacaftor on physical activity and exercise tolerance in three adults with Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A15

Background
The combination of the cystic fibrosis transmembrane conductance regulator (CFTR) corrector lumacaftor with the potentiator ivacaftor has been approved for the treatment of patients with cystic fibrosis (CF) homozygous for the Phe508del CFTR mutation. The major benefits of lumacaftor-ivacaftor therapy may reside in the reduction of pulmonary exacerbations and in a lower annual loss of lung function. However, there are no reports detailing the effect of lumacaftor-ivacaftor on daily physical activity (PA) and exercise tolerance.

Materials and methods
We performed incremental cardiopulmonary exercise testing (CPET) and we assessed PA pre- and post 2 years initiation of lumacaftor-ivacaftor in three adults with CF (CFTR genotype F508del/F508del and chronic airflow obstruction. CPET-related measurements included oxygen uptake (VO2), carbon dioxide production (VCO2), ventilatory profile, heart rate (HR) and oxygen pulse (V02/HR) throughout exercise and at lactic threshold (LT) and peak. LT measures represent sub-maximal exercise related data. PA was assessed using the accelerometer Sense Wear Pro3 Armband.

Results
An improvement of daily physical activity following lumacaftor-ivacaftor was accompanied by clinically significant improvements in exercise tolerance. Specifically, mild (>3.2 metabolic equivalents (METS)) PA improved by +13% in patient 1, +84% in patients 2 and +89% in patient 3; time spent in PA increased by +59% in patient 2 and +115% in patient 3; active energy expenditure improved +54% in patient 2 and +142% in patient 3. VO2 uptake expressed as percentage of predicted significantly increased both at LT and at peak (patient 1 +19 and +33, patient 2 +17 and +42, patient 3 +40 and +20, respectively).

Conclusions
Daily physical activity levels and exercise tolerance improved after two years of LUM/IVA therapy, despite no patient education or lifestyle advice. All patients provided informed written consent for this case report.

A16
Effectiveness of Ivacaftor in Cystic Fibrosis Patients with Gating Mutations: a Two-Years Single Center Experience
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Background
Ivacaftor is a cystic fibrosis transmembrane conductance regulator (CFTR) potentiator, approved for patients with CF with gating and residual function CFTR mutations. We report the results of a single-center observational study investigating its effects in CF patients with gating mutations.

Materials and methods
Patients with gating mutations were recruited to an open-label study evaluating Ivacaftor. Primary outcomes included: lung function, sweat chloride, sputum microbiology, number of pulmonary exacerbations and weight gain.

Results
7 subjects (6 females) were enrolled and completed 24 months follow-up on Ivacaftor; mean age at enrollment was 29.3 years (range 12 – 45 years) with 1 subject <18. Two subjects were excluded from the analysis because of very low adherence to the treatment, documented by large variations of sweat Cl and missed collect of Ivacaftor at the pharmacy. Another patient was excluded because not able to perform the lung function tests, having very severe lung disease with Burkholderia cepacia infection. The remaining 4 participants experienced significant improvements in ppFEV1 (mean absolute increase of 18% at 1 month that was sustained along the follow up period (24 months); sweat chloride decreased from a mean value of 101 mmol/l at baseline to 39 mmol/l at 24 months. Mean absolute BMI increased of 2.6% after 1 month of treatment until 10.8 % at 12 and 24 months. Pulmonary exacerbations were absent and no subject needed hospitalization during the follow up. Microbiology of sputum was unchanged. No safety concern was reported.

Conclusions
Patients with gating mutations experienced improved lung function and nutritional status, and a very pronounced decrease of pulmonary exacerbations. This study supports ongoing use of Ivacaftor for patients with these mutations. Despite the promising nature of Ivacaftor, our data suggest that adherence rates can be suboptimal. Monitoring and motivational interventions are important also for this treatment.

A17
Effects of Ivacaftor/Lumacaftor in Cystic Fibrosis Patients Homozygous for F508del CFTR Mutation
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A17

Background
The combination of a CFTR corrector (lumacaftor) and a CFTR potentiator (ivacaftor) is the first CFTR targeted drug approved for clinical use (Orkambi) for patients affected by cystic fibrosis (CF) homozygous for F508del. In placebo-controlled clinical trials (TRAFFIC/TRANSPORT) the mean absolute improvement in the percentage of predicted forced expiratory volume in the 1st sec (FEV1) ranged from 2.6 to 4.0% and the rate of hospitalization or use of intravenous antibiotics was lower in the group treated with Orkambi [1].

Materials and methods
We describe the clinical and functional effects on CFTR of this drug in 44 CF patients treated at the CF Center of Verona for at least 3 months: prescriptions and follow up were performed according to the rules of Agenzia Italiana del Farmaco (AIFA). Effects of Orkambi were investigated on nutritional status (weight; Body Mass Index, BMI), CFTR function (sweat chloride by Gibson and Cooke), lung function (FEV1), frequency of pulmonary exacerbations (PE, European Consensus Group definition, Bilton et al., 2011) during the first year of treatment. The duration of therapy ranged from 91 days to 4.8 years.

Results
We observed a mean of absolute FEV1 increase of 2.6%, + 8.4 SD, a mean BMI increase of 0.3, ±1 SD, a mean weight increase of 1.1 kg, ±2.6 SD, a mean reduction of 0.800 +0.941SD of PE requiring IV antibiotics or hospitalization, a decrease of sweat chloride values of 19.2 + 12.5SD vs the mean value of the year before the treatment. In 4 patients the treatment was suspended because of adverse effects, most frequently thoracic oppression. In males and females we obtained very similar results. In severe patients (FEV1 < 50%) we obtained lower improvement of nutritional status than in the others (p<0.05) while other effects were not significantly different in these two groups.

Conclusions
CF patients treated with Orkambi at the CF Center of Verona had better nutritional status, lung function and less PE than before treatment, consistently with their improved CFTR function. Although conditions in clinical use are different than in clinical trials the effects on relevant clinical outcomes have been achieved at levels thereabout expected according to the results obtained in the clinical trials. The variability of response to this drug among homozygous F508del CF patients reported in this study suggest the relevance of personalized medicine and new drug development for CF.

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A18
Effect of Ivacaftor on comorbidities in a patient with Cystic Fibrosis (CF) and pancreatic sufficiency
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A18

Background
Generally patients with Cystic Fibrosis (CF) with pancreatic sufficiency have an increased risk to experience acute recurrent pancreatitis. This occurrence is likely due to diminished ductal flow in pancreatic ducts, with inflammation events and inaprt activation of pancreatic digestive enzymes. This risk is related to CFTR dysfunction, causing a lack of fluid secretion. News are emerging on the potential effect of Ivacaftor on comorbidities in CF [1]. We describe a case report about a patient with CF and acute recurrent pancreatitis in treatment with Ivacaftor.

Case report
Giada is a 10 years girl, affected by CF (CFTR genotype: G1244E/3849 +10kbC>T) with chronic lung disease, bronchiectasis, chronic infection by MRSA, Pseudomonas Aeruginosa intermittent airway colonization and pancreatic sufficiency. From April 2017 she is in treatment with Ivacaftor. Since 2 years old she had 7 episodes of acute pancreatitis characterized by severe abdominal pain and increase of serum lipase and amylase about 4XULN. All episodes required hospitalization and treatment of symptoms according to standardized protocol. During episodes other causes of acute pancreatitis were ruled-out. Endoscopic retrograde cholangiopancreatography was performed to exclude bilo-pancreatic malformation. Between the acute events serum pancreatic enzymes remained upper normal values. Already after 8 weeks of Ivacaftor treatment we observed a normalization of both amylase and lipase values that persistently were in the normal range during the 18 months follow-up. The patient has never had pancreatitis. Sweat test decreased from 72 mmol/l to 27 mmol/l. Lung function tests persist in the normal range. Long term follow-up could confirm the effect of Ivacaftor on other comorbidities.

Conclusion
Our experience confirms preliminary results about the positive role of Ivacaftor on controlling recurrence of acute pancreatitis in CF. We suggest that the potentiator could ameliorate pancreatic fluid secretion reducing the risk of pancreatic injury.

Patient’s parents gave consent for the publication of clinical data.

Reference
Carion A, Drucy S, Steven D, ET AL. Reduction of Recurrence Risk of Pancreatitis in Cystic Fibrosis With Ivacaftor: Case Series. J Pediatr Gastroenterol Nutr 2018;66:451-453.
A19 Real-life effectiveness of Lumacaftor/Ivacaftor in Cystic Fibrosis: a single center experience
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A19

Background
Lumacaftor/Ivacaftor (Orkambi, Vertex Pharmaceuticals) is the first CFTR modulator approved for patients with cystic fibrosis (CF) homozygous for the CFTR F508del mutation. This combination therapy is able to increase the amount of functioning CFTR protein on the cell surface. We report the experience at our center with patients on therapy with Orkambi.

Material and methods
We evaluated patients treated with Orkambi since 2016. Patients performed visits after 4 weeks from baseline and then every 3 months. Sweat Cl, BMI, vital signs, physical examination, renal and hepatic profile, spirometry, six-minutes walking test (6MWT), CFQR, sputum culture were evaluated at each visit. All adverse events were registered.

Results
17 patients, mean age at baseline 25.4 years (range 15.3-43.2), 3 in the pediatric age (15.9 years, range 15.3-16.9), 6 F, were studied. 17/17 had pancreatic insufficiency, 7/17 CF-related diabetes, 1/17 CF-related liver disease (Child Pugh A), 1/17 was on oxygen therapy. Mean follow-up period was 16.6 months (range 5-24.2). At baseline: mean sweat Cl value 102.5 mEq/l (range 91-111); mean BMI 21 kg/m2 (17.5-26.3); mean ppFEV1 67 (range 19-110) and ppMMEF 36 (range 5-111); mean 6MWT 597 mt (range 415-740); mean CFQR respiratory score 67.8 (range 27.7-88.9). Through the follow-up period, mean sweat Cl decreased of 13% (range 9-17); mean BMI increased of 10.6 (range 1-21); mean 6MWT were observed. Pulmonary exacerbations were dramatically reduced in 12/17 patients. Respiratory domain of the CFQ-R score improved with a mean gain of 7 (range 0-22.2) from the baseline, at 1 year. Microbiology was unchanged. Cytolysis and cholestasis lab tests improved in the patient with liver disease. 3/7 patients with diabetes decreased their insulin need. Only 1 patient discontinued treatment after 15 months, because of lung transplant. No safety concerns were reported. Adherence to the treatment was good.

Conclusions
The relevant decrease of pulmonary exacerbations rate has been confirmed as the main efficacy measure targeted by Lumacaftor/Ivacaftor. Lung function tests were unchanged during therapy. CFQR-score improvement is probably related to the decrease of pulmonary exacerbation. Use of Lumacaftor/Ivacaftor was safe, even in subjects with severe lung disease and liver disease. Further long term real-life studies, on large number of patients, are needed to evaluate the effectiveness and the prognostic impact of Orkambi.

A20 Lumacaftor/Ivacaftor: use in a 12-year-old patient with Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A20

Cystic fibrosis (CF) is a life-limiting disease that is caused by defective or deficient cystic fibrosis transmembrane conductance regulator (CFTR) protein activity. Phe508del is the most common CFTR mutation. CF affects approximately 70,000 people around the globe including more than 30,000 in Europe. CF is a multisystem disease affecting organs where CFTR is expressed (airways, gastrointestinal and the reproductive tract). Lumacaftor is a CFTR corrector that has been shown to correct Phe508del CFTR misprocessing and increase the amount of cell surface-localized protein. Ivacaftor is CFTR potentiator that increases the open probability of CFTR channels.

Case report
M. P., male 12 years old. Diagnosis of CF in complete form with meconium ileus. Sweat test: 96 mEq/L Cl-. Genetics: F508del/F508del. Intermittent colonization by Pseudomonas aeruginosa from the first months of life. Poor nutritional situation with slow weight gain. In 2015, diagnosis of GH deficiency, starting from January 2016 GH treatment. In August 2017, in consideration of the “critical need”-status, he started taking the pediatric dosage of Lumacaftor/Ivacaftor as it was part of the compassionate use program. For the first week he took Lumacaftor/Ivacaftor 100/125mg every 12h preceded by short acting bronchodilator. From the second week began full pediatric dosage 200/250mg every 12h. At the age of 12 years of age he continued with the same dosage until the time the marketing of the association Lumacaftor/Ivacaftor was made available for patients over 12 years of age. During one year of observation the patient presented an improvement of FEV1 equal to 14% of the starting value, and of the FVC equal to 30%; the 6MWT performance showed an increase of the distance traveled of 285 meters measured at the T12. The assessment of the nutritional status of the child showed a slight increase in weight and height, despite the concomitant use of GH treatment, with a consequent increase in BMI of +0.6. During the treatment, no respiratory exacerbations were observed compared to at least 4 episodes that occurred during the previous 12 months. By submitting the patient to the CFQ-R quality of life questionnaire, the score for respiratory symptoms improved significantly from 50 to 83. The sweat test also showed a reduction in chloride concentration. The only parameter that has not undergone changes has been the microbiological data because the patient has maintained colonization by Pseudomonas aeruginosa.

Conclusion
The use of this therapeutic strategies gave new perspective of life to the patients who carried a Phe508del mutation in CFTR. Parents gave consent to patient data publication.

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A21 Alternative Therapies in patients with Cystic Fibrosis who carry minimal residual function mutations
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A21

Background
Waiting for all available new modulators could be approved for patients with Cystic Fibrosis (CF) who carry class I and/or class II mutations not homozygous for F508del mutation alternative therapies could be evaluated. A previous phase-2 clinical trial, showed that oral cysteamine (CYS), already approved for the treatment of cystinosin, combined with the kinase-inhibiting epigallocatechin-gallate (EGCG,
a safe over-the-counter approved nutraceutical), could be effective in terms of clinical response and functional rescue of CFTR [1]. On the basis of these results we used as off-label therapy this combined treatment for two selected patients.

Case report 1
Matteo (genotype: F508del/2183AA>G) is a 7 years boy who presented chronic lung disease, bronchiectasis, liver steatosis, kidney tubular dysfunction and pancreatic insufficiency. One year treatment with CYS (30 mg/Kg qds) plus EGCG (270 mg qd) was completed. During one year follow-up he improved his lung function: FEV1% increased from 92 to 113%, while MMEF% from 64 to 78%. The sweat chloride decreased from 117 mmol/l at baseline to 55 mmol/l after 12 months. No pulmonary exacerbation was registered compared to one exacerbation in the previous year. The patient reports an improvement of daily exercise capacity.

Case report 2
Ivana (genotype: N1303K/L1065P) is an 18 years girl affected by a severe lung disease, atelectasis, chronic pancreatitis, liver disease treated with UDCA, frequent respiratory exacerbations requiring hospitalization, multidrug allergy, CF related diabetes and pancreatic insufficiency. She assumed the combined therapy with CYS plus EGCG through 2 years. During two years follow-up she improved lung function: FEV1% increased from 68.7 to 79.9%, while MMEF% from 22.3 to 32.6%. The sweat chloride decreased from 113 mmol/l at baseline to 55 mmol/l after 24 months. The transmembrane conductance flow (measured by SPG on nasal brushing sample) gained 50% of wild type after 2 years of treatment. Pulmonary exacerbations reduced in severity with a reduction in the number of hospitalizations of 50% (from 6 to 3) during 2 years follow-up.

Conclusion
Our experience confirms previous data of efficacy and safety of the association CYS plus EGCG in a selected group of patients. In order to confirm the efficacy of CYS plus EGCG combination in subjects with CF carrying two minimal residual function CFTR mutations, a Phase III randomized-controlled multicentre study has to be encouraged.

Patient's parents gave consent for the publication of clinical data.

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GI/NUTRITION

A22
Treatment of Distal Intestinal Obstruction Syndrome (DIOS) in cystic fibrosis: proposal of a multicenter protocol
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A23
Lung clearance index and 24 h pH-metry in Cystic Fibrosis: which link?
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Background
Gastroesophageal reflux (GER) is common in patients with cystic fibrosis (CF) and is often regarded as playing a role in the pathogenesis of CF lung disease. Individuals with CF have many predisposing factors to the development of GER, with a reported prevalence ranging from 35 to 81%. Both the acid and nonacid components of GER may have an effect on lung disease. The lung clearance index (LCI) is a measure of ventilation distribution obtained from the multiple-breath washout (MBW) test. As the MBW test uses relaxed tidal breathing without the requirement of an maximal effort, it is an attractive test for infants and pre-school-aged children. LCI is abnormal in a significant proportion of pre-school children with cystic fibrosis (CF), and correlates with the presence and extent of structural lung disease detected via chest computed tomography (CT) and lower respiratory tract inflammation and infection.

Materials and methods
We enrolled a cohort of 14 patients with CF both children and adults who were investigated with 24-h pH-metry. All patients were free from proton pump inhibitors (PPI) therapy for 1 month and in a stable phase of the disease. At the same time (one day before or after the exam) we performed the study of Lung Clearance Index (LCI) with Multiple Breath Washout (MBW) technique.

Results
We found the presence of relevant episodes of acid gastroesophageal reflux in 10 (71%) patients (6 adults and 4 children). Of these patients, all the 6 adults and only 1 child out of 4 had higher LCI values than normal. Among pediatric subjects with normal 24-h pH-metry 2 presented normal values of LCI and 2 abnormal.

Conclusions
In general, 71% of subjects present acid gastroesophageal reflux confirming that GER is common in patients with CF. In detail, all adults (100%) seem to show a link between abnormal LCI and 24-h pH-metry. Differently, only half (50%) of children showed abnormal normal 24-h pH-metry and only 1 of them showed an abnormal LCI. According to our preliminary data, now we are performing
24-h pH-impedance test in children to establish the weight of basic reflux in children with CF. Further studies are needed to validate implications for Lung disease in patient with Gastro-eposophageal Reflux in CF.

A24
Intestinal inflammation may correlate with ultrasound abnormalities in Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A24

Background
Inflammation in patients with Cystic Fibrosis (CF) is not limited to the respiratory tract, but plays an important role also in the gastrointestinal tract. Fecal calprotectin (FC) is a marker of intestinal inflammation, however its use in CF is controversial. Recent studies support the use of intestinal ultrasound to assess the degree of inflammation in chronic bowel disease.

we have investigated the relationship between high FC levels and the intestinal wall thickening, measured by ultrasound examination, or clinical manifestations in CF patients.

Materials and methods
Data collected for the analyses include age, gender, CFTR mutations, sonographically measured colonic wall thickness, FC levels measured by ELISA, pancreatic insufficiency status (defined as fecal elastase value < 200 μg/g stool), FEV1 as a marker of lung function.

Results
40 CF patients were consecutively enrolled (22 males; mean age 9.6 ±4.2 years; range 3-17 years); 33/40 were pancreatic insufficient and 6/40 had also severe liver disease. FC values were over the cut-off in 16/40 patients, and the intestinal wall thickening abnormal in 15/40 patients, an increased appendicular thickness in 3/40 patients, and an increased wall intestinal thickening in 10/40 patients, an increased appendicular thickness and 10/40 had increased FC value in absence of intestinal abnormalities. Among 7/40 patients with pancreatic insufficient only 2 showed an increase of intestinal wall. FEV1 was available in 31/40 CF patients and 10/30 showed values between 40% and 80%. Only 3/10 patients showed an increased wall intestinal thickening and nobody had increased FC. Among 6 patients with severe liver disease 3 showed an increased wall thickening of which only one has elevated FC. No correlation was found between pathological wall intestinal thickening and/or FC and clinical data such as recurrent abdominal pain, meconium ileus at diagnosis and distal intestinal obstruction syndrome.

Conclusions
Our results show a lower percentage of patients with wall intestinal thickening than data previously reported. In patients with CF, FC could not be a marker of intestinal inflammation as reported in chronic inflammatory bowel disease. Our data show 38% of enrolled patients with elevated FC in disagree with literature [1-2]. Currently preliminary data are not sufficient to reinforce the hypothesis of a correlation between inflammatory marker and wall thickness in children with CF.

References
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2. Parisi GF, Papale M, Rotolo N, et al. Severe disease in Cystic Fibrosis and fecal calprotectin levels. Immunobiology 2017, 222: 582-586

A25
Two cases of Cystic Fibrosis onset with acute pancreatitis and a slow resolution
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A25

Case report 1
Ilary, 7 years, comes to our observation for acute slow-resolution pancreatitis. Second-born from normal course of pregnancy, emission of meconium occurred within the first 24 hours of life. Normal stature-weight growth. During the hospitalization a positive result of the sweat test was carried out and consequently the genetic study was performed with a compound heterozygosis of the CFTR gene (F508del / T388I mutation); the latter is associated with a slightly expressed phenotype, especially with regard to the pancreas secretory function.

Case report 2
Giulia, 11 years old, anything relevant in remote medical history, regular weight-weighted growth. She arrives in emergency area for abdominal pain in the epigastric site radiated to the back, with increasing intensity, not responsive to antalgic therapy. At the blood chemistry tests, high values of pancreatic enzymes are found. Excluding infectious and malformative causes of acute pancreatitis, in the suspicion of cystic fibrosis (CF), a sweat test resulted positive, and genetic testing showed mutation in heterozygosity of G542X and 711 +3A G intron 5. The latter mutation intervenes on the splicing of CFTR and allows the synthesis of a partially functional protein, or a reduced portion of normally functioning CFTR.

In both cases other exams such as broncho-aspirate culture, chymotrypsin and faecal elastase were in normal and it was undertaken fasting, infusion support and subsequently therapy with pancreatic extracts and ursodeoxycholic acid, with gradual normalization of enzyme levels.

Conclusion
Pancreatitis is a relatively rare complication affecting 1-2% of CF patients and may be an initial isolated manifestation of the disease and therefore be the first indicator for the diagnosis of CF. This complication occurs in cases with a functioning pancreas, so in order to manifest inflammation, it is necessary that the organ is still able to secrete enzymes. Approximately 10% of CF patients maintain good pancreatic function until adulthood or for life; they have at least one mutation of the CFTR gene defines as mild. In some of these mild forms, the pancreas causes partial stasis of secretion in the pancreatic canalliculi, inducing inflammation with important abdominal pain and marked increase in pancreatic enzymes in the serum. Follow-up is important for understanding the possible onset of pancreatic insufficiency. In the presence of pancreatitis history, without triggering causes, both in a child and in an adult, it is advisable to investigate for CF.

Parents gave consent to patient data publication.

A26
Elastography as a diagnostic tool to assess liver disease in Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A26

Case report 1
T338I mutation); the latter is associated with a slightly expressed phenotype, especially with regard to the pancreas secretory function.

Case report 2
Giulia, 11 years old, anything relevant in remote medical history, regular weight-weighted growth. She arrives in emergency area for abdominal pain in the epigastric site radiated to the back, with increasing intensity, not responsive to antalgic therapy. At the blood chemistry tests, high values of pancreatic enzymes are found. Excluding infectious and malformative causes of acute pancreatitis, in the suspicion of cystic fibrosis (CF), a sweat test resulted positive, and genetic testing showed mutation in heterozygosity of G542X and 711 +3A G intron 5. The latter mutation intervenes on the splicing of CFTR and allows the synthesis of a partially functional protein, or a reduced portion of normally functioning CFTR.

In both cases other exams such as broncho-aspirate culture, chymotrypsin and faecal elastase were in normal and it was undertaken fasting, infusion support and subsequently therapy with pancreatic extracts and ursodeoxycholic acid, with gradual normalization of enzyme levels.

Conclusion
Pancreatitis is a relatively rare complication affecting 1-2% of CF patients and may be an initial isolated manifestation of the disease and therefore be the first indicator for the diagnosis of CF. This complication occurs in cases with a functioning pancreas, so in order to manifest inflammation, it is necessary that the organ is still able to secrete enzymes. Approximately 10% of CF patients maintain good pancreatic function until adulthood or for life; they have at least one mutation of the CFTR gene defines as mild. In some of these mild forms, the pancreas causes partial stasis of secretion in the pancreatic canalliculi, inducing inflammation with important abdominal pain and marked increase in pancreatic enzymes in the serum. Follow-up is important for understanding the possible onset of pancreatic insufficiency. In the presence of pancreatitis history, without triggering causes, both in a child and in an adult, it is advisable to investigate for CF.

Parents gave consent to patient data publication.
Background
Cystic Fibrosis (CF) is an autosomal recessive genetic disease. CF-associated Liver Disease (CFLD) is a common complication and represents the third cause of death in 2.5% of patients with CF. In clinical practice liver biopsy is not routinely applied to evaluate the severity of liver disease in CF, as it is a painful and invasive procedure. Since much more studies support the use of liver ultrasound to correctly assess the degree of hepatic involvement, we aimed to study a group of children diagnosed as having CF to evaluate the correlation between liver stiffness assessed by ARFI elastography and conventional US qualitative examination.

Material and methods
Data were collected from 34 consecutively enrolled CF patients including age, gender, anthropometric measures, CFTR mutations. Liver stiffness derived from acoustically generated tissue shear wave propagation (Shear-wave elastography-SWE) and Ultra Sound (US) parameters of liver pathology such as hepatomegaly, variation in echogenicity, fibrosis, signs of portal hypertension were also registered in all patients in a bladder manner.

Results
Preliminary results include data from 34 CF patients (17 Males; mean age 8.9 years, range 3.3-15.6). All of them had pancreatic insufficiency and carried minimal residual function mutations on both alleles. 10/34 (29.4%) (8 Males; mean age 12.3 range 7.9-15.0) patients showed hepatic stiffness over the cut-off value (range 6.4-15.0) compared to normal value < 6.3 kPa. In 8/10 patients liver stiffness alterations were concordant with US signs of hepatic abnormalities (steatosis, fibrosis), while 2/10 didn’t show any other sign of liver pathology. In 24/34 patients with liver stiffness in the normal range 6 patients showed US qualitative signs of mild liver abnormalities as steatosis and one with fibrosis.

Conclusions
According to previous data our preliminary results show a correlation between conventional US and increased values of liver stiffness, examined by Shear-Wave elastography [1] in order to detect CFLD. More data are needed to identify liver stiffness as a diagnostic tool of liver disease with advantage of real-time imaging and lower cost and to correlate it to clinical characteristics.

Reference
1. Aqul A1, Jonas MM, Harney S, Raza R, et al. Correlation of Transient Elastography With Severity of Cystic Fibrosis-related Liver Disease. J Pediatr Gastroenterol Nutr. 2017; 64: 505-511.

Table 1 (abstract A27). Clinical characteristics

| CFTR GENOTYPE | DATE OF BIRTH | FECAL ELASTASE 1 VALUE mcg/g of feces (sampling date) | FECAL FATS (normal value <3%) | AGE AT STOP (year) | PERT (year) |
|---------------|---------------|------------------------------------------------------|-------------------------------|-------------------|-------------|
| F508del/2789+5G>A | 22/06/2012 | 30 (10/7/12) 17% | Fecal Elastase 1 | 1 | 1 |
| F508del/2789+5G>A | 07/09/2006 | 169 (April 2007) | <3% | 8 |
| F508del/2789+5G>A | 07/09/2006 | 64 (20/12/07) | 20-33% | - |
| F508del/2789+5G>A | 07/09/2006 | 495 (27/8/13) | <3% | - |
| F508del/2789+5G>A | 07/09/2006 | 500 (8/10/13) | <3% | - |
| F508del/2789+5G>A | 07/09/2006 | 471 (31/12/13) | 5-12% | - |
| F508del/2789+5G>A | 07/09/2006 | 500 (30/04/14) | <3% | - |
| F508del/2789+5G>A | 07/09/2006 | >500 (28/9/15) | - | - |
| F508del/2789+5G>A | 07/09/2006 | 82,82 (23/06/14) | 21-10-10% | 4 |
| F508del/2789+5G>A | 07/09/2006 | >500 (22/02/18) | - | - |
| F508del/2789+5G>A | 07/09/2006 | >500 (22/03/18) | - | - |
| F508del/2789+5G>A | 07/09/2006 | >500 (27/04/18) | - | - |

A27
Transitory Pancreatic Insufficiency in mild form of Cystic Fibrosis: the importance of follow-up with Fecal Elastase-1
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A27

Background
About 85-90% of Cystic Fibrosis (CF) patients has pancreatic insufficiency (PI). PI involves the CF patients with two CFTR “severe” mutations (functional class I, II, III), while the presence of at least one copy of an allele of class IV-V correlates in the majority of cases with a pancreatic sufficiency (PS) phenotype. Although genotype predicts the development of PI there is a clinical variability due to the interaction of other genes (e.g. inflammatory response genes) and environmental factors. The loss of pancreatic function generally appears during the first months and years of life, thus pancreatic function is an important clinical marker of the progression in CF. In order to treat the malabsorption improving the nutritional status it is important to determine the pancreatic functional status since the first month of life in screened babies. In clinical practice the most common pancreatic function test used is measurement of Fecal Elastase-1 (E1) as a good marker of pancreatic status. The test had a 93% sensitivity and a 93% specificity with a <200 ug/g feces cut off. Normally E1 reaches adult levels in the first two weeks of age, thus this method is useful for screening PI/PS in screened infants.

Cases report
Here we present 4 cases of CF females carrying at least one "mild" mutation of CFTR gene. In the first months of life, they had E1 insufficient levels (Table 1), then infants were labeled as PI and pancreatic enzyme replacement therapy (PERT) was quickly begun. Monitoring pancreatic status during clinical follow-up, we assisted to a spontaneous increase of E1 levels, that reached stably PS values allowing us to stop PERT and improve their quality of life.

Conclusions
Genotype predicts the maintainance of exocrine PS in CF, as the presence of at least one CFTR mutation of functional class IV-V generally correlates with PS, but there is a clinical variability determining fluctuation of E1 values. We reported four "mild" CF cases initially labeled as PI in the first months of life, however during the follow-up we assisted to a spontaneous raise of E1 values reaching PS range as expected from their genotype. Monitoring regularly pancreatic status also in PI children who carry CFTR mutations with a residual function is necessary to avoid unnecessary PERT.

The families of the patients have given written consent for the publication of the data.
A28
Percutaneous Endoscopic Gastrostomy (PEG): the Experience of The Regional Reference Center for Cystic Fibrosis of Milan
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A28

Background
Malnutrition is a complication of Cystic Fibrosis (CF) and has been recognized as a negative prognostic factor. Therefore malnutrition has to be corrected and a step wise approach has been recommended including enteral nutrition (EN) via percutaneously placed gastrostomy (PEG) tube [1]. The aim of this study was to investigate the efficacy of EN via PEG to improve nutritional status, respiratory function and incidence of complications.

Materials and methods
A retrospective cohort study on CF patients who underwent PEG between 2000 and 2016 was performed at our Center. Sixteen patients were included in the study with a median age of 13 years, 94% with pancreatic insufficiency. The anthropometric data were recorded 6 months prior PEG placement, at the time of placement and 6 months after. Patients were classified according to CF Foundation (CF) nutritional criteria. Malnutrition was considered to be present in patients with BMI <10th percentile up to 19 years and <18.5 kg/m² over 20 years. In addition, in 15 parents of 13 patients, the level of parental stress was investigated through a self-administered questionnaire; results of PEG-group (n=4) were compared to a group of control subjects with a CF child but not needing EN.

Results
Malnutrition was present in almost all patients who underwent PEG, with the exception of 3 children in whom PEG was necessary due to total aversion to oral feeding. After 6 months of EN median values of the Z-score of W/L or BMI increases by 1 standard deviation. At the start of EN, 13 patients (81.3%) were malnourished; 6 months after the malnourished subjects decreased to 31.3% (N=5).

Complications (mechanical, gastroenterological, metabolic) occurred in 50% of patients and were solved; only in one case NE was stopped. The mean percentile value of the Total Stress levels falls within the norm in both groups. Only one mother reported a value in the range of clinical interest due to the coexistence of other problems in addition to CF.

Conclusions
Patients needing PEG constitute a small minority in CF population; from our experience it emerges that EN via PEG is safe and helps to achieve an adequate weight when usual interventions fail. Patients with PEG must be evaluated from a nutritional point of view at every clinical check and supported in the management of this device.

Reference
1. Smyth AR, Bell SC, Bojcin S, et al. European Cystic Fibrosis Society Standards of Care: Best Practice guidelines. J Cyst Fibros. 2014;13: S23–S42.

A29
Nutritional status in the first 2 years of life of children with Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A29

Background
Optimizing growth from early childhood in Cystic Fibrosis (CF) is critical, as lung function and survival correlate strongly with nutritional status. Guidelines recommend a persistent optimal nutritional status. The objective of this preliminary study is to evaluate nutritional status in a cohort of children with CF in the first 2 years of life, in follow-up at Cystic Fibrosis Centre of Naples

Materials and methods
Data from 12 children (5 Males) affected by CF, born between 2015 and 2016, were retrospectively evaluated. CF was diagnosed in 10/12 by neonatal screening and in 2/10 for suggestive symptoms but with negative newborn neonatal screening. 9/12 were pancreatic insufficiency (PI). We evaluated data about nutritional status (body weight, height and P/A ratio with the relative percentile) according to CDC growth curves, feeding modalities, prescriptions of nutritional supplements, clinical outcome and vitamin D levels. Furthermore, we investigated mothers anxiety, evaluated by STAI -Y questionnaire, in relation to nutritional status of enrolled children.

Results
Results of anthropometric parameters are summarized in Table 1. 4 patients at a high risk of malnutrition improved through personalized nutritional intervention at 2 years of age. From the analysis of feeding modalities, 4/12 children were breastfed, 5/12 were formula-fed exclusively and 3/12 received mixed feeding. The dietary intake for children was about 120% (range 100%-130%) estimated average requirement. Insufficient children received PERT at a dosage increasing from 2500 lipase/g fat units at the diagnosis to 3300 lipase/g fat units at 24 months of follow-up in order to correct malabsorption. In particular, steatocrit values normalized on average from 34% (range 12%-40%) to 2.32% (1% - 6.5%) All enrolled children received salt supplementation. Despite vitamin supplementation in all children since diagnosis only one with severe pancreatic insufficiency showed hypovitaminosis D.

Beyond their nutritional status preliminary data concerning 8 mothers showed a level of anxiety (trait: 35-63 state: 33-72) measured by STAI -Y questionnaire with no correlation to nutritional status.

Conclusions
Early individualized nutritional intervention is useful to ensure adequate growth and correct nutritional status in CF. More data are needed to correlate nutritional status and mother anxiety that could impact on rescue of malnutrition.

Table 1 (abstract A29), Results

| Lenght | pc At diagnosis | 2 years N=12 | Weight/length | pc At diagnosis N=12 | 2 years N=12 |
|--------|----------------|-------------|--------------|---------------------|-------------|
| <5°p   | 4              | 0           | <5°p         | 0                   | 0           |
| 10°-25°| 0              | 1           | 10°-25°      | 2                   | 2           |
| 25°-50°| 2              | 3           | 25°-50°      | 3                   | 3           |
| 50°-75°| 6              | 5           | 50°-75°      | 6                   | 6           |
| 75°-90°| 0              | 3           | 75°-90°      | 1                   | 1           |
| pc=percentile |

A30
Severe anemia as rare onset clinical sign of Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A30

Case Report
We report the case of a male, only one son of unrelated parents, born at the 37th week by caesarean section, weight at birth of 2900 gr. At the age of 3 months, he came to our attention for suspected cystic fibrosis (CF) due to the positivity of neonatal screening. The
patient appeared very pale and edematous, but in good general conditions and normal vital parameters. We decided to perform a blood count, the sweat test and directly the genetic screening for CF. The examination of the blood count showed a severe normocytic anemia: Hb 5.5 g/dl RBC 2,000,000/mmcc, MCV 67 fl, PLT 364,000/mmcc, WBC 10,430/mmcc. The child was hospitalized, and he underwent other laboratory tests that showed hypoproteinemia (4.7 g/dl), hypoalbuminemia (1.43 g/dl) and increased hemolysis indices (total bilirubin 4.7 mg/dl, direct bilirubin 1.8 mg / dl). The first clinical suspect was a hemolytic anemia, then excluded. He performed parenteral albumin supplementation and blood transfusion with normalization of laboratory values. The sweat test was pathological and the genetic test for CF identified the presence in heterozygosis of two CFTR variants (c.1521-1523delCTT, (delta)F508 e CFTR:c.3846G>A (W1282X) related to CF.

Conclusion
In the literature there are currently only two recent works of Sismanlar T. of 2016 and Aricò M. of 2018 describing the onset of CF with severe anemia and hypoproteinemia, as rare symptoms related to poor intestinal absorption. Parents gave consent to data publication.

INFECTION/ MICROBIOLOGY

A31
Is Serum Amyloid A a marker for diagnosing cystic fibrosis pulmonary exacerbation?
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Background
Identification of pulmonary exacerbation (PE) in CF patients is important for proper treatment. Serological markers as C-reactive protein (CRP) and leucocyte (WBC) counts are not very specific and not included in the Fuchs criteria for PE diagnosis. Serum amyloid A (SAA) protein, are a family of apolipoproteins expressed or secreted by liver during the acute phase of inflammation. SAA genes are regulated by the proinflammatory cytokines (IL-1, IL-6, TNF-α) and induced rapidly under inflammatory conditions like exposure to bacterial LPS. SAA is also involved in amyloidogenesis and tumor pathogenesis. At low concentrations (10-100ng/ml), as in early inflammation, SAA induces chemokines or matrix degrading enzymes and functions as a chemoattractant. When an infectious or inflammatory stimulus persists, the liver continues to produce SAA that becomes a direct bacterial opsonin or interferes with viral infection of host cells. AA amyloidosis is a rare, but severe complication of CF with predominant renal involvement. Aim of the study is to evaluate the usefulness of SAA as a marker of inflammation and a diagnostic criterion for PE.

Materials and methods
Between January 2016 and December 2017 we collected serum SAA, CRP and WBC at the time of admission and at discharge for PE (defined according to the Fuchs criteria) in 22 CF patients. Details of the study population are shown in table 1.

Results
A total of 40 hospital admissions for PE, in 22 patients (M/F: 11/11, mean age at admission: 11.5 years, range 1-21) were analysed. All 40 events met the Fuchs criteria for PE. The mean serum SAA at admission was 166.5 ng/ml (range 2-937 ng/ml, normal value: < 6 ng/ml). The mean serum CRP at admission was 21.65 mg/l (range <2.9-139 mg/l, normal value: < 2.9 mg/l).

16/40 (40%) had a negative CRP at admission while only 4/40 (10%) had a negative SAA (p<0.05). 4/16 (25%) events with negative CRP had also a negative SAA. WBC were in the normal range for age in 36/40 events (90%). At the time of discharge the SAA was negative in 28/40 (70%) events (mean SAA: 9.9 ng/ml, range <6-74 ng/ml).

Conclusions
Literature is still discussing the definition of PE. New and more sensitive biochemical markers of early inflammation are necessary to diagnose PE. SAA may represent a more sensitive biochemical marker than CRP and WBC for PE diagnosis and might be included, after an adequate validation study, in the definition criteria for PE.

Table 1 (abstract A31). Characteristics of the study population

| Patients (n°) | 22 |
| Sex (M/F) | 11/11 |
| Pulmonary exacerbation (n°) | 40 |
| Pulmonary exacerbation/patients | 1.8 |
| Pancreatic sufficiency (n°) | 1 (4.5%) |
| MRSA chronic pulmonary infection (n°) | 4 (18%) |
| PA chronic pulmonary infection (n°) | 13 (59%) |
| Mean age at admission (years) | 11.47 (1-21) |
| Mean duration of hospitalization (days) | 15 ±2 |
| Mean serum SAA (ng/ml) (range) | 166.5 (2-937) |
| Mean serum CRP (mg/l) (range) | 21.65 (<2.9-139) |

Table 45 (Suppl 1):A32

A32
Palivizumab prophylaxis to prevent respiratory syncytial virus infection in infants with Cystic Fibrosis (CF). Experience of the last five years
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A32

Background
Respiratory syncytial virus (RSV) is associated with worsening respiratory symptoms in children affected with CF increasing also the hospital admission. Palivizumab is the only preventive therapy, since no vaccine against VRS is now available. No conclusive data are available regarding the use of Palivizumab in CF population because few studies have been carried out.

Material and methods
All patients diagnosed with CF and in follow up to our Unit born between 2014 and 2018 were included in a retrospective study. All data on hospital admissions caused by respiratory disease were collected until September 20, 2018. RSV testing was performed from nasopharyngeal aspirates. All patients received Palivizumab in the first year of life. The aim of our study was to obtain epidemiologic data on the incidence of RSV infection in children with CF less than 24 months of life who received Palivizumab.

Results
We identified a cohort of 36 patients in follow up in our Unit (F 22, M 14) with CF born between 2014 and 2018 with a diagnosis by screening or symptoms before 24 months of life, who received Palivizumab ( five doses, 15 mg/kg for dose) in the first year of life. 31 of them had a diagnosis for screening and 5 for symptoms. Two patient with CF diagnosed for symptoms are awaiting vaccination in the next October due to a recent diagnosis. Three patients showed a RVS infection during admission in hospital for respiratory symptoms, so the incidence of RSV infection in our cohort is 8.8%. One patient born at 28+3 weeks of gestational age, affected with meconium ileus with...
Ceftolozane/tazobactam for the treatment of MDR Pseudomonas aeruginosa in a patient suffering for Cystic Fibrosis waiting for lung transplant

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A34

Background

Cystic Fibrosis (CF) is an autosomal recessive disease due to the mutation of the gene that encodes the Cystic Fibrosis Transmembrane Regulator (CFTR), which regulates and is involved in the transport of electrolytes. Incidence: 2.000-3.000 new cases a year. CF is a multisystem disease affecting organs where CFTR is expressed (airways, gastrointestinal and the reproductive tract). The main pathogenic germs are Staphylococcus aureus (Sa), Pseudomonas aeruginosa (Pa). In CF a targeted and aggressive therapy of chronic Pa infections is one of the mainly strategy to allow the improvement of the prognosis. The new combination of β-lactam and a β-lactamase inhibitor such as Ceftolozane-Tazobactam shows a rapidly bactericidal action, a linear kinetics, it spreads well in the lung and acts on numerous mechanisms of resistance of Pa.

Case report

Female aged 16, weighing 51 Kg, suffering from CF in a complete form, waiting for lung transplants, is hospitalized for fever and dyspnoea. EGA and blood exams showed a septic state. She arrived with Levofloxacin, Tobramycin, Meropenem antibiotic iv treatment. Cultural examinations were performed and started antibiotic therapy with Ceftolozane/Tazobactam and Phosphomycin. From the fifth day of therapy the clinical conditions gradually improve and also the blood tests showed a reduction in CRP. Therapy is continued through 20 days. In the meantime, a subsequent sputum culture examination isolated 2 strains of Pseudomonas (mucoid and wrinkled) with a sensitivity similar to that of the entrance except for a resumption of sensitivity to Piperacillin-Tazobactam. So we decided to modify the therapy from ceftriaxone to Piperacillin/Tazobactam and Colistina iv. After 7 days the clinical conditions worsened, with a reappearance of fever and increase in inflammation indices. The last sputum culture test document the reappearance of resistance to Piperacillin-Tazobactam, therefore we decided to stop the therapy and to resume the association Phosphomycin+ Ceftolozane/Tazobactam. There was a progressive improvement of the clinical conditions.

Conclusion

The patient came to our observation in a highly compromised general state, that is, with an exacerbation of her chronic Pa MDR infection. We started a combination therapy with the aim of improving the clinical outcome i waiting for the lung transplant. The choice fell on Ceftolozano-Tazobactam due to its strong anti-pseudomonas action, used with good clinical evidence also in rescue therapies. We did not observe any side effects and the
hepato-renal and medullary parameters remained in the normal range. Parents gave authorization to patient data publication.

A35
Longitudinal metagenomic 16s sequencing of a CF infant’s respiratory samples as a tool for microbiome monitoring

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Introduction
Pulmonary environment plays a major role in modulating CF lung disease, thus, accurate identification of the whole respiratory microbiome has a crucial role for the management of patients, the improvements of their healthcare and quality of life. CF infants’ microbiome analyses can give a better understanding on the associations between bacterial populations and patient conditions during their lives, starting from the very beginning, and can better show how this relationship affects pulmonary microenvironment and the evolution of CF disease. Sequencing analyses and comparison with cultural approaches were performed on 10 clinical samples that cover the entire life of a single 2-years patient to obtain a longitudinal study on the variation of the respiratory microbiome compared to the culture-based methods.

Materials and methods
From January 2015, ten samples of nasopharyngeal aspirates were collected from a single patient starting from the first microbiological exam of his life. These samples firstly underwent to microbial analysis performed by routine cultural standards for CF samples, stored by frozen, and, retrospectively, were processed by New Generation Sequencing (NGS). Briefly bacterial DNA was extracted using Qiangen kit and 16s library preparation protocol was adjusted for poor quality samples (as infants’ ones are). Raw reads taxa were assigned by BLAST and visualized by MEGAN software.

Results
NGS demonstrated how commensal flora undergoes to several modifications during sampling period, underlying the presence of different families never discriminated by microbial cultures such as anaerobic population (Prevotellaceae and Fusobacteriaceae) that were quite abundant (up to 18%) into the microbiome. Also, variations of different families were observed over time and prevalence of Staphylococcus aureus, considered by culture the main pathogen in this patient, was sometimes different if compared with families identified by NGS.

Conclusions
Nowadays, the culture-based methodology is considered the gold standard for microbiological diagnostics: although it has high specificity, this method has low sensibility due to several problems, such as the difficult isolation and detection of pathogens from the polymicrobial flora, the presence of unculturable pathogens and anaerobes that cannot be detected using standard culture-based conditions and the presence of pathogens extremely adapted to the pulmonary microenvironment that grow after several days (up to 7 days) under peculiar conditions (growth at 30°C or in micro-aerobic atmosphere).

For all these reasons, culture-based methodology cannot characterize all the different bacterial populations in each sample and so culture independent technologies, such as NGS technology, are needed to overcome the previous issues.

A36
The role of nasal washes in patients with Cystic Fibrosis affected by chronic rhinosinusitis
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Background
Cystic Fibrosis (CF) presents multiorgan manifestations that include chronic rhinosinusitis (CRS) with or without nasal polyposis. Nasal washes (NWs) are widely used in clinical practice especially in CF patients, although their effectiveness on Ear Nose Throat (ENT) symptoms is controversial.

In this study we evaluate the performance and the safety of a NWs solution, with or without surfactant, to reduce symptoms and bacterial load.

Materials and methods
We enrolled 20 CF patients (mean age: 27.6 years) with CRS, confirmed by nasal endoscopy. All patients, colonized by Pseudomonas Aeruginosa, performed daily a NW by physiological solution or by saline solution with surfactant (Naridek). All patients, at the time of enrollment, filled out a sinonasal questionnaire (SANOQ11) and they received instructions for proper washing. During follow-up, we evaluated the reduction of the bacterial load in the nasal lavage. We assessed the nasal cavities by endoscopy (2.7 mm 30° rigid endoscope - Storz, Tuttinglen, Germany) according to a modified Lund Kennedy endoscopic scoring system:

- rhinorrhea (present = 0, mild = 1, purulent = 2);
- edema, and hyperemia (absent = 0, mild = 1 or severe =2);
- nasal mucosa (eutrophic = 1; hyperemic = 2; dystrophic = 3);
- left and right turbinate hypertrophy (none = 0; mild = 1; medium = 2; and serious = 3).

All subjects underwent the Sniffin’Sticks to evaluate the olfactory performance.

Results
Twelve patients completed 4 months of treatment: 6 patients performed the treatment with Naridek and 6 patients with physiological solution. Due to the small sample size, the scores were added together to have an overall indication of the treatment (Table 1)

The bacterial colonization in NWs shows no statistically significant difference. However, in 2 patients, we detected a reduction of the bacterial load, while there was no difference in the saline-treated group.

Conclusions
Considering our small sample we can only draw some great deal to think about:

- treatment with NWs allows an improvement of the ENT symptoms and is well tolerated by patients. These data are confirmed by the ENT signs score and by the reduction of the SNAQ11 score in both treatment arms;
- the solution with surfactant (Naridek) significantly improves the ENT signs and decreases the nasal endoscopy and the SNAQ11 scores;
- no benefit was detected at the evaluation of olfactory performance.

In conclusion, even if further confirmations are necessary on broader cases, it seems to emerge as significant the role of surfactant in the therapeutic advantage of NWs.
A37
Burkholderia gladioli: a new potential hazard in Cystic Fibrosis patients?
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A37

Background
Burkholderia gladioli is an opportunistic Gram-negative, not belonging to Burkholderia cepacia complex (BCC). Its clinical impact ranges from transient to chronic infections and it is responsible of poor post-lung transplant outcomes. Therapeutic strategies are not standardized. We describe 5 cases of B. gladioli infection in cystic fibrosis (CF) patients to highlight the wide variability in the clinical course.

Materials and methods
Presumptive identification of B. gladioli was assessed by MALDI-TOF and confirmed by 16S sequencing. The isolates from single patients and stains isolated from persistently infected patients were genotyped both with BOX-PCR approach and MLST analysis.

Results
We genotyped strains from 4 out of 5 patients. The isolates were genetically unrelated. Instead, strains isolated from each chronically infected patient showed an identical profile. MLST profiles showed the environmental origin of the bacteria.

Conclusions
Some B. gladioli may cause transient infections, especially if strains are susceptible to antibiotics. Cases of chronic infections have been reported and antibiotic strategies are not clearly established. In our study genotyping showed that all patients were infected with genetically distinct strains, while a single strain was responsible of persistent infection in two cases. Large multicenter studies are needed to investigate epidemiology, clinical course and effect of B. gladioli on lung disease in CF.

Written informed consent for publication was received from the patient/parent of the patient

A38
Ceftazidime–Avibactam and Ceftolozane/Tazobactam: new antimicrobials against multidrug-resistant Pseudomonas aeruginosa in Cystic Fibrosis
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A38

Background
The emergence of multidrug-resistant Pseudomonas aeruginosa (MDR-PA) strains is an important and growing issue in the care of cystic fibrosis (CF) patients. Ceftazidime–avibactam and ceftolozane/tazobactam are novel antimicrobials combining a third-generation cephalosporin with β-lactamase inhibitor the use of which are restricted to some clinical cases, including CF patients infected with MDR-PA. In this study, we assessed the ceftazidime–avibactam and ceftolozane/tazobactam susceptibility of MDR-PA isolated from adults with CF.

Materials and methods
MDR-PA (strains resistant to all antibiotics in two or more classes of anti-pseudomonal agents), were isolated from CF patients in our center in the last year. The antimicrobial susceptibility for all the isolates were performed by using the micro broth dilution method following EUCAST guidelines.

Results
The activity of ceftazidime/avibactam and ceftolozane/tazobactam was assessed against 29 MDR-PA isolates collected from 15 CF patients (mean age 34.0±11.0 yrs) chronically infected. We also analysed 32 strains from a control group of 12 CF patients (mean age 29.7±9.8yrs) with chronic no-MDR Pseudomonas aeruginosa infection. The rate of susceptibility of MDR isolates to colistin and ceftazidime/avibactam were 100% and 48.3% respectively, followed by 27.6% to ceftolozane/tazobactam (MIC50 16/32 mg/liter) and tobramycin. Ceftazidime-avibactam (MIC50/90, 4/16 mg/liter) was the most active (highest susceptibility rates) compounds after colistin. All other antibiotics were active only in less than 10% of the isolates, no isolates were susceptible to ciprofloxacin. In the non-MDR isolates the susceptibility rates were 96.9% and 90.6% to ceftolozane/tazobactam (MIC50/90, 1/2 mg/liter) and ceftazidime-avibactam (MIC50/90, 2/8 mg/liter) respectively. All antimicrobials resistance rates are shown in the table 1.

Conclusions
Ceftazidime–avibactam and ceftolozane/tazobactam are novel antimicrobial drugs targeting difficult to treat Gram-negative organisms. The data showed that these antimicrobials were highly active in vitro against CF isolates of Pseudomonas aeruginosa, including isolates that exhibit resistance to multiple drug classes. This study demonstrates excellent susceptibility profiles specially for ceftazidime–avibactam against MDR Pseudomonas aeruginosa strains collected from the sputum of individuals with CF.

| Table 1 (abstract A38) | Pa antimicrobials resistance rates |
|------------------------|----------------------------------|
| GEN | AMI | TOB | CIP | CAZ | AZT | IMI | MEM | TZP | CZA | C/T | COL |
| Pa | 3.7 | 3.4 | 27.6 | 0.0 | 10.3 | 6.9 | 10.3 | 17.2 | 13.8 | 48.3 | 27.6 | 100.0 |
| MDR | % | S | | | | | | | | | |
| Pa no | 65.6 | 75.0 | 87.5 | 21.9 | 81.3 | 31.3 | 59.4 | 31.3 | 15.6 | 90.6 | 96.9 | 100.0 |
| MDR | % | S | | | | | | | | | |

Abbreviations: Pa, Pseudomonas aeruginosa; MDR, Multidrug-resistant; GEN, Gentamicin; AMI, Amikacin; TOB, Tobramycin; CIP, Ciprofloxacin; CAZ, Ceftazidime; AZT, Aztreonam; IMI, Imipenem; MEM, Meropenem; TZP, Piperacillin / tazobactam constant 4; CZA, Ceftazidime/avibactam; C/T, Ceftolozane/tazobactam 4; COL, Colistin

Table 1 (abstract A38). Pa antimicrobials resistance rates
RESPIRATORY

A39
Cystic Fibrosis Sinus Score for paranasal sinuses complications of Cystic Fibrosis: a three-years long experience

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A39

Background
Management of paranasal sinuses complications in Cystic Fibrosis (CF) is a challenge for the otolaryngologist. Paranasal sinuses system represents a P. aeruginosa reservoir affecting lower airways system. Early diagnosis and follow-up of ENT complications of CF, like mucoceles and polyposis, are important to treat them. A clinical-symptoms, radiological score for treatment, stratification and evaluation of CF patients has been developed.

Materials and methods
Between May 2015 and May 2018, 52 patients (27 males, 25 females; mean age 15.90 years, range 6-30 years) attending our ENT-CF clinic were evaluated for paranasal sinuses complications with maxillofacial CT; cone-beam CT (CBCT) investigation was used for pediatric patients. Radiological results were described qualitatively and quantitatively using Lund-Mackay score. CT and CBCT report with the radiological score, endoscopic Meltzer’s Score and SNAQ-11 questionnaire for clinical ENT symptom were used for the assessment of a CF sinus score (CFSS), with a maximum score of 112 points. ENT treatment was chosen depending on CFSS results.

Results
CT and CBCT guided Lund-Mackay scores: mean result 20.25 (range 6-24). By CFSS findings, 20 patients (38%) were addressed to endoscopic sinus surgery (mean result 67.5; range 32-80); the remaining patients were addressed to follow-up/medical therapy (mean result 23.8; range 12-42). Most relevant radiological findings were pluri-sinonasal disease, medialization of maxillary sinuses walls, mucoceles, and bone resorption. At endoscopic evaluation polyposis was the most frequent finding (61.5%); finally, nasal obstruction and rhinorrhea were registered as the most referred symptoms (42% and 21% each).

Conclusions
Our CFSS results confirmed that CF patients present an early clinical objectivity of nasal polyposis with pluri-sinonasal disease, asymptomatic in most of the cases. CBCT has demonstrated to be a suitable radiological investigation and follow-up for CF patients, due to its reduced radiation exposition. Future objectives of CFSS is to identify CF patients who can benefit the most from surgery and to become a mean for comparison among ENT specialists treating CF.

Went to our referred center complaining dyspnea at rest, worsening of airway clearance and the need for more oxygen support during the day.

A40
High flow nasal cannula for exacerbation of dyspnea in patient with Cystic Fibrosis: a case report

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A40

Background
We presented the case of a 27-year-old man with Cystic Fibrosis (CF) due to a DF508 homozygotic mutation. The patient is colonized with Pseudomonas aeruginosa, Aspergillus fumigatus and Mycobacterium acccessus that were treated in the past with no evidence of reactionation, and an allergic bronchopulmonary aspergillus (ABPA) was detected. He is currently treated with a CFTR corrector (lumacaftor) and a CFTR potentiator (ivacaftor). Patient was treated with low oxygen flow (LOF) support continuously throughout the day. Before the admission, the health perception was assessed according to CFQ-R Teen/Adult Italian Version 2.0. The patient went to our referred center complaining dyspnea at rest, worsening of airway clearance and the need for more oxygen support during the day.

Materials and methods
Patient was admitted with high respiratory rate (35 breaths/min). His first gas assessment showed hypoxemia (pO2 55.8 mmHg) and low pO2/FiO2 ratio (1.36). Chest X-ray was suggestive for a new opacity. Then, high flow nasal therapy (HFNC) (AIRVO 2 Fisher and Paykel) was administered to the patient (34 °C, 50 L/min and FiO2 42%). Patient continued high flow nasal cannula therapy almost continuously for seven days and for a further four nights. We prescribed HFNC domiciliary at night time.

Results
Patient reported great comfort and showed high tolerance to this device and setting. Respiratory rate decreased shortly after the start of high oxygen flow therapy. Gas assessment at the end of treatment with HFNC revealed increased level of pO2 and pO2/FiO2 (pO2 85.9 mmHg, pO2/FiO2 2.45). The patient acknowledged an improvement in cleaning the secretions. At the time of discharge, the patient pointed out the great benefits of the HFNC treatment and his general health perception increased according to CFQ-R Teen/Adult Italian Version 2.0.

Conclusions
In patients with CF the clearance of airway secretions remains one of the most common issues. It is well known that HFNC delivers warm and humidified gas and thus improves mucociliary function by maintaining clearance to a greater degree than other methods for delivering oxygen. Due to the capacity of the device to reach higher and relatively fixed FiO2 than LOF, hypoxemia improves throughout the treatment. Furthermore, HFNC generates positive end-expiratory pressure which reduces the work of breathing and respiratory rate. The low burden of the device allows the patient to continue the treatment also at home with potential benefits in terms of health perception, cleaning secretions and respiratory physiotherapy. The patient gave the consent to publish clinical data.

A41
Observational study of sleep respiratory breathing disorders and preventive noninvasive ventilation in cystic fibrosis: preliminary results and proposal of longitudinal and multicenter study

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A41

Background
In Cystic Fibrosis (CF) lung disease is characterized by progressive airflow obstruction, due to mucus plugging and airway inflammation, causing destruction of the lung parenchyma secondary to bronchiec-tasis. These alterations result clinically in an increase of the work of breathing, leading to alveolar hypoventilation predominantly during sleep, exercise and acute respiratory exacerbations. In clinical practice, nocturnal cardiorespiratory polygraphy (PG) role is growing and may represent an important diagnostic test compared with conventional pulmonary function test as spirometry and lung clearance index (LCI). Furthermore, signs of lung function decline, marked principally by impaired gas exchange and/or increased work of breathing during sleep, may be represent powerful and validated criteria to start preventive nocturnal noninvasive ventilation (NPPV) in CF patients. The purpose of this study was to identify respiratory patterns over the spectrum of disease severity in patient with CF. The overall
hypothesis for the current study is that patients with CF demonstrate gas exchange abnormalities and increased respiratory loads during sleep, reported and recognized by conventional cardiorespiratory PG and also, using non-invasive ventilation as preventing respiratory failure in contrast with conventional practice in clinical studies.

Materials and methods
Analysis of nocturnal breathing patterns and gas exchange on PG was performed in patients with CF (30) patients with different clinical characteristics. They performed also spirometry and LCI with multiple-breath washout (MBW) test.

Results
Children with CF demonstrated lower oxyhemoglobin saturation (95% ± 1.6%), higher respiratory rate (20.5 ± 6.9 breaths per minute). The Apnea Hypopnea index differ between age, particularly in preschool age (5.5 ± 2.7 events per hour.). Ventilation distribution outcomes (LCL) was significantly higher (>7) in patients with advanced CF. 5 patients with serious nocturnal hypoxemia and/or increasing respiratory loading started nocturnal noninvasive Bi-level ventilation. After 6 months, control polygraphy showed an improvement of respiratory pattern.

Conclusions
NPPV may be represent a potential treatment in CF to prevent lung function decline, in contrast with conventional use in acute respiratory failure, or as a bridge to pulmonary transplant. Using Bi-level PPV, upper airway obstruction and/or work of breathing induced by intrinsic positive pressure of end expiration (PEEP) are prevented by expiratory positive airway pressure (EPAP) and thus pressure support (PS) can be triggered easily by the patient. In conclusion, different ventilatory modes have been used in patients with CF, but bi-level positive pressure ventilation is used by the majority of the patients for the comfort and the security regarding to the inspiratory pressures.

ECMO as a bridge to recovery from severe exacerbation in non-end stage lung disease of a patient with Cystic Fibrosis: a case report

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A42

Background
Extracorporeal life supports (ELS) are devices that enable direct oxygenation or CO2 extraction from the blood. In recent years their use has increased among CF patients with end-stage respiratory condition and failure as a bridge for lung transplantation (LTx). Nevertheless, extracorporeal membrane oxygenation ECMO support (ECMO) is considered when a patient presents a rapid deterioration of a chronic lung disease and only when it has been already included on a LTx waiting list or an evaluation process has already started.

Case Report
A 25 years old woman with CF, CF-related diabetes (CFRD), CF liver disease (CFLD) with portal hypertension and a history of moderate but unstable lung disease, chronic Pseudomonas Aeruginosa colonization and allergic bronchopulmonary Aspergillosis (ABPA) presented with fever (38°C), persistent cough, excessive stagnation of thick mucus and dyspnea. Chest X-ray showed diffuse bilateral par-enchymal thickening, several bronchiectasis with mucoid impaction, presence of venous-sub pleural bubbles. Intravenous antibiotics and corticosteroids were started along with intensive regimen of respiratory physiotherapy for airway clearance. During the first hours her status deteriorated, so that she needed to be transferred to the intensive care unit (ICU) where she required inotropic support mode. After 4 hours, despite IMV at maximal protective pressure and maximal antibiotic coverage, arterial blood gas (ABG) analysis showed a persistent unresponsive hypoxemia, severe respiratory acidosis and severe hypercapnia (pH 7.18, pCO2 87 cm H2O), so veno-venous ECMO was initiated. After 40 hours, ABG analysis improved, so she was successfully extubated and placed on continuous NIV. During ECMO period, mucolytic therapy was doubled and two sessions per day of airway clearance were performed by respiratory physiotherapists. Ventilation parameters improved, so she has been “bridged” from NIV to heated humidified high-flow nasal cannula therapy. ECMO de-cannulation occurred on day 10 and after 48 hours she was discharged from ICU on spontaneous breathing with 5 lt/min oxygen supplementation. Respiratory physiotherapy was continued, reassessed and optimized along with a training program. She was discharged home on after 10 days without oxygen supplementation.

Conclusion
Considering clinical history and the stage of pulmonary disease, this patient was not considered yet for LTx. ECMO is an important support to severe CF exacerbations unresponsive to conventional treatments in patients that are not yet candidates to LTx. ECMO “bridge-to-recover” use is emerging in CF and could extend native lungs function in order to take time for waiting list evaluation and optimize allocation.

The patients gave the consent to publish clinical data.

Table 1 (abstract A42). Arterial Blood Gas analysis, Mechanical Ventilation and ECMO parameters during the ICU hospitalization

| Time from admission | pH | pO2 [mmHg] | pCO2 [mmHg] | SaO2 [%] | FiO2 [%] | IPAP [cmH2O] | EPAP [cmH2O] | HFNC Flow [L/min] | Blood Flow [L/min] | Gas Flow [L/min] |
|---------------------|----|------------|-------------|---------|---------|--------------|--------------|-----------------|----------------|----------------|
| 0 hr                |    |            |             |         |         |              |              |                 |                 |                |
| 1 hr                | 7.18 | 141.0 | 87.9 | 98.8 | 100 | 28 | 8 |
| 4 hr                | 7.40 | 95.7 | 50.0 | 98.3 | 25 | 25110 | 8 | 2.92 | 2 |
| 10 hr               | 7.44 | 68.7 | 45.0 | 94.3 | 30 | 23110 | 8 | 2.9 | 2.5 |
| 38 hr               | 7.50 | 77.5 | 45.6 | 96.2 | 35 | 22116 | 8 | 3.47 | 3 |
| 41 hr               | 7.51 | 87.0 | 44.7 | 97.6 | 42 | 10 | 7 | 3 | 3 |
| 48 hr               | 7.52 | 64.7 | 42.7 | 94.3 | 40 | 12 | 7 | 3.56 | 5 |
| 60 hr               | 7.55 | 88.2 | 36.8 | 98.0 | 40 | 14 | 6 | 3.8 | 5 |
| 99 hr               | 7.42 | 88.0 | 47.4 | 96.7 | 60 | 12 | 7 | 40 | 2.8 | 5 |
| 110 hr              | 7.51 | 68.8 | 34.7 | 94.3 | 60 | 40 | 2.73 | 7 |
| 120 hr              | 7.50 | 64.6 | 35.0 | 93.0 | 50 | 30 | 2.54 | 6 |
| 162 hr              | 7.47 | 74.2 | 34.3 | 95.0 | 50 | 30 | 2.9 | 2.5 |
| 182 hr              | 7.47 | 100.0 | 34.4 | 98.2 | 45 | 30 | 2.26 | 4 |
| 191 hr              | 7.47 | 94.7 | 35.8 | 97.6 | 46 | 30 | 1.8 | 2.5 |
| 207 hr              | 7.44 | 10.0 | 37.2 | 94.0 | 50 | 20 | 1.87 | 2.5 |
| 232 hr              | 7.44 | 88.7 | 40.8 | 96.8 | 45 | 20 | 1.98 | 2.5 |
| 231 hr              | 7.45 | 105.0 | 36.6 | 98.2 | 60 | 20 | 2.37 | 3.5 |

Removing...
PHYSICAL AND RESPIRATORY THERAPY

A43

Comparing the efficacy of two different training modality in adult patients with cystic fibrosis: a retrospective observational study

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A43

Background

It is well established that exercise has multiple benefits for people with cystic fibrosis (CF), but the evidence for health effects of specific sport disciplines is poor. Therefore, we retrospectively examined the impact of two different training modality - jumping on a trampoline and cycling on a stationary bike - on spirometry, exercise capacity measured with 6 Minute Walk Test (6MWT), exertional dyspnea and the perception of muscular fatigue.

Materials and methods

A six months retrospective observational study was conducted from 1st March to 31st August 2018 at the Cystic Fibrosis Centre, Policlinico Umberto I, Rome, Italy. Data were collected from medical records of 30 adult CF patients (19 F; mean age 31.1 ± 10.2 yrs) hospitalized for pulmonary exacerbation (mean hospital stay 14.2 ± 5.5 days). The main outcomes were spirometric values (FVC, FEV1, FEF25-75), secondary outcomes were 6 minutes walked distance, dyspnea (measured with a Borg scale before and after the 6MWT) and muscular fatigue perception measured in the same way. A comparative analysis was conducted to test the significance of differences within same group and between groups. SPSS version 24.0 was used for data entry and statistical analysis. Univariate comparisons were made using Student’s t-test. Alpha was set at 0.05.

Results

Participants of both groups had improvement in all outcomes, presumably due to the overall treatment received. The study shows no statistically significant difference between groups for the main outcome while the 6 minutes walked distance (from 566.1 ± 47.0 m to 604.7 ± 43.8 m; p=0.001), dyspnea (from 6.5 ± 1.5 to 3.7 ± 1.3; p=0.004) and muscular fatigue perception (from 3.8 ± 1.5 to 1.7 ± 0.7; p=0.001) have improved in the trampoline group in a statistically meaningful way. The retrospective design of the study did not allow to know patients’ preference between jumping and cycling.

Conclusions

Jumping on a trampoline could be a valid training modality for people with CF, but more research is needed to better understand its efficacy, safety and patients preference compared with other training modality.

A44

Manual therapy for musculoskeletal pain in cystic fibrosis: review of the literature

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A44

Background

Pain is a common complication in patients with cystic fibrosis (CF), negatively associated with pulmonary exacerbations, quality of life and treatment adherence. Moreover, with the increased life expectancy the prevalence of musculoskeletal complications is growing, and the use of physiotherapy techniques to minimise these problems has increased. The aim of this study is to review the efficacy of physiotherapeutic manual techniques to manage musculoskeletal pain in patients with CF.

Materials and methods

MEDLINE, CINAHL, PsychINFO, Scopus, CFDDB and trials registries via WHO Portal were searched since 2000 up to September 14t 2018. Search terms included (cystic fibrosis) AND (musculoskeletal pain) AND (manual therapy) AND (sham therapy OR rest OR exercise OR drug OR surgery).

Results

364 studies were identified, of which 4 (190 participants) met the inclusion criteria: 3 randomised controlled studies (85 participants) and one non-randomized study (105 participants). The study size ranged from 20 to 105 participants (age 18 or older). Duration of treatment in the included studies ranged from one day to six months. Due to data heterogeneity no meta-analysis could be performed. Manual therapy (in the form of osteopathic manipulative treatment, spinal joint and intercostal mobilization, soft tissue therapy, massage or chiropractic) was compared with sham therapy or conventional care (no manual therapy). Three RCT compared manual therapy with sham therapy or no treatment (usual care). Primary endpoints were FVC, and a composite outcome reflecting the intensity of pain and the number of painful days over the previous month. Secondary endpoints were self-assessment of breathing, pain and anxiety level measured with a questionnaire, quality of life (Cystic Fibrosis Questionnaire), need for analgesics. The studies showed no statistically significant difference between the treatment groups and the control groups, it is however interesting to note that in one study 26 patients (81%) indicated in a questionnaire that they had been interested in the study and 15 of 16 patients (94%) in the treatment arm were very satisfied with the treatment received. One non-randomized trial studied the effect of a combination of musculoskeletal physiotherapy techniques and massage therapy (a single treatment session) on musculoskeletal pain and ease of breathing (VAS) showing a significant reduction in both outcomes, irrespective of clinical status (acute versus stable patients).

Conclusions

Larger and more powerful studies are needed to determine whether manual therapy techniques could be a valuable therapeutic option in the management of musculoskeletal pain for patients with cystic fibrosis. Patients interest and satisfaction also encourage further research.

A45

Intrapulmonary percussive ventilation with high-frequency chest wall oscillation: make way for positive vibration? A case report

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A45

Background

Six-year-old girl admitted in the Cystic Fibrosis Centre, “Policlinico Umberto F” of Rome, for a pulmonary exacerbation colonized with Pseudomonas Aeruginosa. She had a CT thorax: various bronchiectasis filled with mucus in lingular area, in superior right lobe and superior left lobe. During hospitalization she received antibiotic therapy, corticosteroids, doxase alpha and, in the first days, daily respiratory physiotherapy with pep-mask. Before the admission, the patient used to have dance classes, four times a week and jumped on a trampoline every day for 45 minutes. The patient’s adherence was good - her family actively participated in treatment plan - but her attitude towards the cure, physiotherapy in particular, was critical, because the girl perceived the therapeutic burden as very high.

Materials and methods

In an attempt to increase efficacy and the girl appreciation for treatment, during the 14 days hospitalization she received the prescribed
A46 Tracking adherence through I-Neb: you never stop learning!
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A46

Background
Time requirements for multiple daily nebulizer treatments are important impediments to the quality of life for most patients with cystic fibrosis (CF). The I-neb Adaptive Aerosol Delivery (AAD) results in shorter treatment times and better deposition of the drug system, if used correctly. It can be used with 2 modality of breathing: the Target Inhalation Mode (TIM) or Tidal Breathing Mode (TBM). Most importantly, it allows patients, care-giver and CF team to have feedback on the ongoing aerosol therapy. The aim of this study was to evaluate the need of an educational intervention (EI), assessed by recorded adherence, inhalation technique and mesh performance on I-neb device.

Materials and methods
The platform Insight Online (IoL) used by respiratory physiotherapists at the CF Centre of Milan was initially screened to detect patients actively on I-neb. Adherence, inhalation and nebulization time, and cleaning data were reviewed and analysed.

Results
After the initial screening, some data were dropped because withdrawal (n=22), patient changed CF center (n=1) and early termination (n=14). There were 43 patients actively on inhaled antibiotics treatment with I-neb on IoL. To date, 30 patients with CF aged 5-34 yrs were considered for the analysis (13 have not been followed-up yet); 27 used TIM and 3 the TBM breathing modality. Overall, I-neb compliance was 72.4 (36.1)% (range 0 to 132%); the mean rest time was 37.7 (31.4)% (range -6 to 88%) and the mesh performance was 78.2 (40.3)% (range 0 to 147%). The mean nebulization time when using TIM was 25’ (23’’ (range 0’ to 81’’); if TBM was used, the mean time was 80’(4’5’’) (range 4’6’’ to 13’0’’). Mean inhalation time per breath using TIM was 2.8’(2.7’) (range 0’’ to 7.9’’), using TBM was 1’0’(0’9’) (range 0.4’’ to 2.1’’).

Conclusions
IoL gives a unique opportunity to monitor patients’ adherence, the performance of the aerosol device and can provides information that could have been missed if using self-reporting. On average, the majority of patients showed good results, however there is a great variability when using this device. Therefore, it may be necessary to provide an EI to train patients at inhaling. It may be also useful to make the training software available at home to improve the inhalation technique.

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A47 Evaluating physical performance in paediatric patients: experience of the Cystic Fibrosis (CF) centre of Milan
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A47

Background
Standardized exercise is part of the regular assessment of patients with CF [1]. It is used for the evaluation of physical limitations, to document exercise-associated symptoms or to screen for possible adverse effects of exercise [2]. Exercise is an important therapeutic modality for individuals with CF, with higher physical activity levels being associated with greater aerobic fitness [3], pulmonary function, glycemic control [4], and bone mineral density. The Godfrey protocol has been used for the clinical assessment of adverse reactions to exercise in CF. The aim of this study is to evaluate exercise tolerance in paediatric patients regularly followed-up at Milan CF centre.

Materials and methods
After reviewing the literature and the feasibility of different exercise testing at our centre, we adopted the Godfrey cycle protocol since November 2014 as routinary assessment of exercise. Informed consent and test operator procedure were then realized. After March 2018, following the donation of a paediatric cycloergometer, we have been testing the paediatric population below 120 cm-height as well.

Results
So far, 63 patients (31 female) with CF have performed the Godfrey test. Mean (SD) age was 13.8 (2) years (range 7.8 to 15.9 yrs) with a FEV1 of 85 (26.2)%pred. 4 patients (6.3%) had FEV1 <40%pred, BMI perc 39.39’(26.5); any tests have been performed with oxygen therapy. HR peak was 171.72 (14.8) beats/min, HR percentage was 88.3% (7.7), Wmax was 122(42.8) watt. Only 35 tests (55.5%) were maximal: 15 for Wmax, 27 for BORG CR10 ≥ 7, 9 for HR peak. Only 7.5% (n=12) of the whole sample showed a normal response to exercise. The average time needed for the test was 100’8” (6’23” to 14’25”).

Conclusion
These data suggest that in the pediatric population exercise testing offers an opportunity of detecting significant abnormalities. It would be important to perform cardio-pulmonary exercise testing in patients with abnormal response.

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QUALITY IMPROVEMENT

A48

The Italian External Quality Assessment. Scheme for Sweat Test: the importance of a good communication with Laboratories

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Background

Sweat chloride is the gold standard test for the diagnosis of cystic fibrosis (CF). Internal and External quality controls are mandatory to assess precision and accuracy of test results and are recommended to improve the performance of the laboratory.

Materials and methods

In 2014 the National Centre for Rare Diseases (CNMR) of the Istituto Superiore di Sanità (ISS) established the Italian pilot external quality assessment scheme for CF sweat test (IEQA-ST). In 2015 this activity was recognized as a third party service carried out by the ISS. The program is performed once a year and participation is voluntary. Private and public laboratories, performing sweat test, are invited to participate. The IEQA-ST consists of a coordination board (managing the working plan) and of an evaluation board (assessing the performance of each laboratory) composed by experts from CNMR-SSS and Scientific Societies (SIFC, SIMMESN and SIBiOC). An IEQA-ST web-based utility is used to facilitate communication and data sharing among all parties. Each round consisted of the following items: 1) meeting between the boards to ameliorate items and criteria to be adopted in the upcoming round; 2) agreement by participants on the general rules and the assessment criteria; 3) delivery of 3 sweat like samples (including mock identification data, clinical and technical information; 4) collection of technical, institutional and analytical results (including report) from participants; 5) assessment of results; 6) a decisional meeting between the boards; 7) delivery of the final report to all labs; 8) annual meetings. The latest item represents the qualifying event of this plan. In 2016 the poor performance criteria was also adopted.

Results

The number of labs involved in this activity increased (from 10 to 16); analytical errors decreased by about 45% from 2016 ahead (2015: 11; 2016: 6; 2017: 6); clinical sensitivity (mainly wrong interpretation of the analytical result) errors increased (2015: 12; 2016: 9; 2017: 13); poor performance labs increased (2016: 3; 2017: 4). Main points of discussion during the annual meeting with all labs included: assessment criteria; final report; mock clinical information. Main feedbacks from labs: to maintain the annual meeting; to improve the content of the final report; to improve the communicating process.

Conclusions

The IEQA-ST scheme is continuously evolving in accordance with national and international sweat test recommendations and feedbacks from participants. In this respect, laboratories opinion on the communication process, the technical support and the discussion about the assessment criteria are the driving force to improve the quality of the scheme.

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EPIDEMIOLOGY AND NEWBORN SCREENING

A49

Hypochloraemic Metabolic Alkalosis: insidious diagnosis

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Italian Journal of Pediatrics 2019, 45(Suppl 1):A49

Background

Cystic fibrosis (CF) is the most common recessive genetic disorder in Caucasians. The suspicion of disease comes from the detection of typical symptoms, familiar history and newborn screening (NBS). Case reports

We describe cases of two children who came to our observation for dehydration and metabolic alkalosis. NBS for Cystic Fibrosis resulted negative.

Clinical Case 1. Male 6 months. No genetic-metabolic diseases in family. History of recurrent ear infections. Some episodes of vomit, fever and hypoxemia have been reported for some days. He appears in poor general conditions, moderate dehydration. EGA: PH: 7.52, HCO3: 36. Na: 129 mEq / L; K: 2.4 mEq / L; Cl: 91 mEq / L. ECG: hypokalemia. Hospitalized in our department he performs urinary electrolytes in the standard, renal function tests within normal limits. Eco-abdomen: negative, not signs of hypertrophic pyloric stenosis. He performs rehydrating therapy with physiological solution and Potassium Chloride with sudden improvement of the general clinical conditions and normalization of the ion and the parameters of the EGA within 24 hours.

Clinical Case 2. Male 10 months. No history of genetic diseases in family. Influenza episode at one month of life, bronchiolitis at the age of 3 months. From 15 days feverish, recurrent vomiting and hypoxemia, reported weight loss (500 g). He comes to our observation for torpor and hyporeactivity. EGA: hypochloremic metabolic alkalosis. He performs urinary electrolytes, renal function tests within normal limits. Eco-abdomen: abdominal effusion, thickening of intestinal loops. Chest x-ray: multiple pulmonary densities. He performs...
rehydrating therapy with physiological solution and Potassium Chloride with sudden improvement of the general clinical conditions and normalization of the ion and the parameters of the EGA within 36 hours. Both children performed screening for infective disease, renal tubular enzymes and Cortisol that were in the normal range. In stable conditions a second test was also performed and it resulted over the cut-off level of 60 mmol/l, therefore diagnosis of CF was performed, subsequently confirmed by genetics (2989+5G>A/4382delA; F508del/PSL, respectively).

**Conclusion**

NBS false negativity can be associated with laboratory errors, bad choice of cut-off values, pancreatic sufficiency, meconium ileus. Considering the high prevalence of false negativity, our centre is working to increase test sensitivity lowering the first IRT cut-off, to make a second test on the same sample (from 54 to 50 ng/ml) and defining the second IRT cut-off 40 ng/ml at 3-4 weeks of age. In the detection of a hypochloremic metabolic alkalosis, in addition to CF, diseases such as RGE, APLV, metabolic diseases, adrenogenital syndrome, hypertrophic pyloric stenosis and tubulopathies must be sought. Patients gave consent to data publication.

### A50

**Cystic Fibrosis Screen Positive Inconclusive Diagnosis (CFSPID): Experience in Tuscany, Italy**

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**Italian Journal of Pediatrics 2019, 45(Suppl 1):A50**

**Background**

The increased implementation of Cystic Fibrosis (CF) newborn screening (NBS) had led to identification of infants with a positive NBS test but inconclusive diagnostic testing, classified as “CF screen positive, inconclusive diagnosis” (CFSPID) in Europe [1–2]. We retrospectively evaluated the prevalence and clinical outcome of CFSPID patients by two NBS programs in the period 2011-2016 at CF Centre of Florence, Italy.

**Materials and methods**

We retrospectively evaluated CFSPID and CF patients diagnosed by CF NBS in the years 2011-2016 and followed until 31.12.2017. In this period, the project was aimed to assess the diagnostic impact of DNA analysis on the NBS algorithm (immunoreactive trypsin (IRT) - meconium lactase – IRT2), IRT was considered elevated for values above the 99th centile laboratory cut-off. The CFSPID definition was according Munck A et al.1 All CFSPID patients repeated SC over 6 months and performed extended CFTR gene analysis (detection rate 98%). During follow up we reclassified children as: CF diagnosis, healthy carrier or healthy in presence respectively of at least two pathological SC or two normal SC for age and 1 or 0 CF-causing mutation. We kept the CFSPID definition when SC was persistent in borderline range or in presence of two CFTR mutations, at least one of which has varying clinical consequence (https://www.cftr2.org/).

**Results**

Of 179684 babies screened from January 2011 to December 2016, 1520 (0.8%) screened IRT-positive value at day 3. Infants called to perform SC test were 359 by NBS algorithm and 181 by DNA analysis. We identified 32 FC diagnosis and 50 CFSPID (CF:CFSPID ratio 0.62:1). One of 179602 (0.0005%) cases resulted false negative by NBS and diagnosed by dehydration with hypochloremic metabolic alkalosis at 11 months. 20/50 (40%) CFSPID cases were diagnosed only by the IRT-DNA algorithm, 13/50 (26%) by only IRT-meconium lactase-IRT2, while both protocols identified the remaining 17 cases (34%). 36/50 CFSPID patients have a conclusive diagnosis on 31.12.2017: 5 (10%) FC, 16 (32%) healthy and 15 (30%) healthy carrier; 14/50 (28%) cases are asymptomatic with persistent borderline SC and followed as CFSPID (CF: CFSPID ratio 2.6:1). CFTR genetic analysis impacts on sensibility and positive predictive value (Table 1).

**Conclusions**

In six years CF: CFSPID ratio modified from 0.64:1 to 2.85:1 and 10% of CFSPID progressed to CF. Genetic analysis improved positive predictive value and identified a higher number of CFSPID infants progressing in CF.

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### A51

**Cystic Fibrosis Screen Positive Inconclusive Diagnosis (CFSPID): nine years of experience in an Italian Cystic Fibrosis Centre (Naples)**

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**Italian Journal of Pediatrics 2019, 45(Suppl 1):A51**

**Background**

Newborn screening (NBS) for Cystic Fibrosis (CF) identifies classic CF and most CFTR-Related Disorder (CFTR-RD), but also equivocal cases not fulfilling the diagnostic criteria for CF. Genetic analysis improved positive predictive value, LR= Likelihood ratio=*

| A50 | CF vs (CFSPID+Healthy) | CF+CFSPID vs healthy |
|-----|-------------------------|----------------------|
|     | IRT-lact-IRT2 | IRT-DNA | IRT-lact-IRT2 | IRT-DNA |
| Sensitivity % | 87 (75-96) | 90 (79-97) | 78 (67-89) | 90 (81-97) |
| Specificity % | 99.84 (99.83-99.86) | 99.92% (99.91-99.93) | 99.85 (99.83-99.87) | 99.93% (99.92-99.94) |
| PPV% | 11 (8-14) | 19* (15-25) | 13 (10-16) | 26* (21-32) |
| NPV% | 99.99% (99.99-99.99) | 99.99% (99.994-99.998) | 99.99% (99.99-99.997) | 99.99% (99.994-99.999) |
| Positive LR | 555 (404-763) | 1140* (829-1567) | 514 (391-677) | 1256* (955-1653) |
| Negative LR | 0.13 (0.10-0.18) | 0.11 (0.08-0.14) | 0.22 (0.16-0.28) | 0.10 (0.07-0.13) |

Positive predictive value; *Negative predictive value; LR= Likelihood ratio;*p<0.001
Materials and methods
From January 2008 to December 2017, at Cystic Fibrosis Center of Naples (Pediatric Unit), 263,479 children were screened; 81 were identified as CFSPID (71%, 44 males) and 33 as CF (29%, 17 males). These patients were monitored to define their evolution through clinical, biochemical, microbiological data and imaging procedure (median follow-up: 3.8 year. Range: 11 months-9.6 years).

Results
By NBS significant differences between CF and CFSPID were detected:
- I IRT: CF (mean ± DS: 140.1 ± 72.3 ng/dl) vs CFSPID (72.5 ± 21.4 ng/dl) p<0.001;
- II IRT: CF (147.7 ± 79.7 ng/dl) vs CFSPID (48.2 ± 10.7 ng/dl) p<0.001;
- ST: CF (77 ± 15.7 mmol/l) vs CFSPID (15.7 ±7.3 mmol/l) p<0.001;
- CF patients had 2 CFTR causing mutations, while among CFSPID, 43 had 1 causing mutation and 1 with unclear phenotypic consequences and 38 had 2 mutations with unclear phenotypic consequences.

The follow-up of patients with CFSPID revealed that:
- these patients had less respiratory symptoms as compared to classic CF;
- none showed pancreatic insufficiency, meconium ileus nor metabolic alkalosis episodes;
- 8 cases developed a clinical feature suggestive of CFTR-RD (mean age: 4.5 years), one patient had a diagnosis of CF (Table 1.)

Conclusions
Our data suggest that:
1. based on our NBS program CFSPID is about 3 folds more frequent than classic CF (CFSPID:CF ratio 2.45:1) (the highest frequency reported so far);
2. few patients may evolve to CFTR-RD/CF;
3. the follow-up of three years suggested by the current literature is not sufficient to identify cases potentially evolving to CFTR-RD/CF;
4. no differences of IRT or ST were detected by NBS program in CFSPID cohort that could predict evolution to CFTR-RD/CF.

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Table 1 (abstract A51). Features of patients with diagnosis of CFTR-RD/CF.

| Patient | Characteristics | Diagnosis | Age of diagnosis | Sweat chloride | CFTR Genotype |
|---------|-----------------|-----------|------------------|----------------|---------------|
| 1       | bronchiectasis  | CFTR-RD   | 8                | 28             | F508de/Q1475X |
| 2       | bronchiectasis  | CFTR-RD   | 8                | 27             | 1717-1G>A/ST12G |
| 3       | recurrent pneumonia, nasal polyp | CFTR-RD | 11 | 7 | F508de/L997F |
| 4       | recurrent pneumonia, chronic sinusitis | CFTR-RD | 3 | 5 | R334Q/L997F |
| 5       | recurrent pneumonia | CFTR-RD | 26 | 5 | F508de/D1152H |
| 6       | recurrent pneumonia | CFTR-RD | 12 | 5 | S1426F/D1152H |
| 7       | recurrent pancreatitis | CFTR-RD | 27 | 5 | L732X/D1152H |
| 8       | recurrent pancreatitis | CFTR-RD | 28 | 5 | F508de/D1152H |
| 9       | sweat positive test | CF | 3 | 62 | c.2657+5G>A/ST13TG |

PSYCHOLOGY

A52

Meaning of “time” after the lung transplantation
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Italian Journal of Pediatrics 2019, 45(Suppl 1):A52

Background
The lung transplant’s experience is a complex situation where a series of factors interact each others. A good physical condition in the post-transplantation period is not always a guarantee of positive emotional perception. The progressive resumption of life, sometimes are not experienced in a good emotional state, as if the patient could not take his “new time”. We report a clinical case of transplant cystic fibrosis patient where the psychological sessions showed this apparent contradiction.

Materials and methods
During the follow-up post-transplantation, we evaluated a patient, currently of 35 years old, with psychological assessment, for a period of six years (2012-2018). To assess depression and anxiety, at first evaluation in 2014, we administered respectively Hospital Anxiety Depression Scale (HADS) and State Trait Anxiety Inventory (STAI), and subsequently, as indicate by the Mental Health International Committee, we administered Patient Health Questionnaire (PHQ-9) and General Anxiety Disorders (GAD-7). Furthermore, during every evaluation, we administered Cystic Fibrosis Questionnaire-Revised (CFQ-R).

Results
During this period the general physical condition of the patient, the lung function and the nutritional state did not show critical aspects. The patient didn’t live any negative personal events. The score of depression, anxiety and quality of life after lung transplant are described in Table 1.

Conclusions
The results of psychological evaluation show higher levels of depression and lower score in QoL than the physical condition could be suggest. This case report underlines that a post-transplant, although without clinical complication, is not always an opportunity for the patients to re-discover all the aspects of the “new life”. This patient, like other similar in our experience, seems not to be able to enjoy his “time” of life. As well as, when the disease was disabling, they do not live their time even after transplantation. These patients are looking for a more intensive life and are worry about death. They have the perception of having lost “time” immeditely, due to cystic fibrosis.

Knowing the average life expectancy in the post-transplant, they live like in a “turned hourglass”, where the transplant is the moment when the countdown started. We should ask ourselves if we are underestimating some deep aspects of their personal experience, consequently limiting the support we can give them. For this reason it is not only necessary do the psychological screening, but also, every time, the clinical interview. In this way we can really know the patient’s needs.

Patient gave the consent for publication.

Table 1 (abstract A52), Patient characteristics

| Diagnosis | 1 year before (2011) | 2 years later (2014) | 5 years later (2017) | 6 years later (2018) |
|-----------|----------------------|---------------------|---------------------|---------------------|
| FEV1      | 22,50%               | 67.10%              | 77.00%              | 79.80%              |
| STAI-1    | /                    | /                   | /                   | /                   |
| STAI-2    | /                    | /                   | /                   | /                   |
| HADS      | /                    | /                   | /                   | /                   |
| PHQ-9     | /                    | 9 (mild)            | 15 (moderate-severe) |
| GAD-7     | /                    | 7 (mild)            | 3 (normal)          |
| CFQ-R     | 70.1                 | 77.6                | 70.4                |
**A53**

**Early Glucose Metabolism Alteration in Cystic Fibrosis Patients is associated with lower lung function**

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**Background**

Cystic fibrosis-related diabetes (CFRD) patients suffer from accelerated rates of pulmonary decline [1], also before the onset of overt CFRD (2-3). This study aimed to determine if in a young CF population early alteration of glucose metabolism correlates with lung function or microbiological respiratory status.

**Materials and methods**

Clinical (BMI and cBMI), functional (FEV1 and FVC % predicted) and microbiological (presence of chronic Pseudomonas infection) data were collected from CF patients (>10 yrs) who underwent a screening oral glucose tolerance test (OGTT) as part of their routine clinical care since September, 2016 to August, 2018. All data were collected at the time of the last OGTT test. Statistical analysis was performed using T-test to compare means, linear and multiple linear regression (Stata/SE 12.0) after normalization for age. A P-value <0.05 was considered significant in all tests.

**Results**

Screening OGTT was performed in 50 CF patients (24M/26F), mean age 17.01 years (± 4.63 yrs), median age 15.84 (range 10.60-31.55). CFRD was diagnosed in 9 (18%), in 11 (22%) an IGT resulted, while 60% (30 subjects) had a normal glucose metabolism. In the table (Table 1) clinical and metabolic data are presented.

**Conclusions**

Fasting blood glucose is not a helpful screening to diagnose early glucose metabolism alteration in CF patients. CF patients with an impaired glucose metabolism (IGT + CFRD) show a statistically significant lower FEV1 and FVC percent predicted. Chronic Pseudomonas colonization was higher in IGT and CFRD patients, however not statistical significant. Our data confirm the need of implementation of annual OGTT test in CF patients >10 yrs to identify early alteration in glucose metabolism, thus prompt nutritional advices and therapeutic measures might be implemented.

**References**

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**ENDOCRINE**

Table 1 (abstract A53). See text for description.

| Characteristics | All (n=50) | NGT (n=30) | IGT (n=11) | CFRD (n=9) |
|-----------------|-----------|-----------|-----------|-----------|
| Age (yrs)       | 17.01 ± 4.63 | 14.76 ± 2.47 | 19.41 ± 5.80 | 20.27 ± 2.86 |
| BMI (kg/m²)     | 20.05 ± 1.63 | 21.54 ± 2.09 | 21.0 ± 1.98 | 19.96 ± 1.64 |
| cBMI            | 40 ± 26 | 40 | 40 | 30 |
| Fasting blood glucose (mg/dl) | 87.05 ± 8.0 | 85.73 ± 5.92 | 89.80 ± 13.09 | 85.33 ± 4.93 |
| 1-h OGTT blood glucose (mg/dl) | 186.35 ± 55.25 | 170.33 ± 39.98 | 216 ± 68.21 | 215.67 ± 85.48 |
| 2-h OGTT blood glucose (mg/dl) | 124.11 ± 40.19 | 104.8 ± 24.06 | 159.1 ± 15.08 | 215.33 ± 120.0 |
| Fasting blood Insulin (μU/ml) | 7.68 ± 10.51 | 9.10 ± 12.08 | 4.7 ± 3.65 | 2 |
| 2-h OGTT blood Insulin (μU/ml) | 56.81 ± 41.04 | 48.69 ± 31.81 | 63.3 ± 44.86 | 74 ± 33.94 |
| FEV1 (% p.)a | 84.20 ± 21.57 | 90.4 ± 19.5 | 85.45 ± 21.33 | 65.88 ± 9.05 |
| FVC (% p.)b | 95 ± 17.20 | 100.77 ± 15.69 | 92.78 ± 18.20 | 80.5 ± 11.13 |
| Chronic Pseudomonas infectionc | 46% | 43.3% | 54.5% | 44.4% |

aFEV1(%): CFRD vs NGT p=0.0015; IGT vs NGT p=0.1; CFRD vs IGT p=0.035.

bFVC CFRD vs NGT p=0.0077

cChronic Pseudomonas Infection X²=0.4, p<0.1 (NS)

**A54**

**Bartter syndrome or Pseudo-Bartter Syndrome in a Cystic Fibrosis infant? One face for two different diseases**

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**Background**

Cystic Fibrosis is a systemic pathology with associated loss of salts with sweat. Pseudo-Bartter syndrome (PBS) is an uncommon cause of metabolic alkalosis that has been described as a presenting feature of CF, as well as a complication in those with known disease. It is accompanied by chronic salt depletion and sometimes failure to thrive without severe dehydration. PBS is characterized by hyponatremic, hypochloremic metabolic alkalosis that mimics Bartter syndrome, but with no pathology in the renal tubules.

**Case report**

We present the case of a 1-year-old male with Eastern Europe origins, affected by CF in homzygous for the F508del. The patient comes to our attention very frequently due to repeated dehydration attacks without severe dehydration. PBS is characterized by hyponatremic, hypochloremic metabolic alkalosis that mimics Bartter syndrome, but with no pathology in the renal tubules.

**Conclusion**

The poor response to therapy and the extremely early onset of such manifestations have made us suspect that the patient may not have the CF-related PBS but the true Bartter syndrome.
Bartter is an autosomal recessive syndrome which manifests with an alteration of the salts reabsorption resulting in extracellular fluid volume reduction with possible low or normal blood pressure. It is caused by mutations of genes encoding proteins that transport ions across renal cells in the thick ascending limb of the nephron also called as the ascending loop of Henle. Specifically, mutations directly or indirectly involving the Na-K-Cl cotransporter are key. Loss of function of this reabsorption system results in decreased sodium, potassium, and chloride reabsorption in the thick ascending limb, as well as abolishment of the lumen-positive voltage, resulting in decreased calcium and magnesium reabsorption. Consequently, Patients with Bartter syndrome may also have elevated renin and aldosterone levels as compensation mechanism. There are 4-5 different type of Bartter syndrome, they are due to compound heterozygous or homozygous mutations in four genes that code for proteins involved in the reabsorption of chloride in the ascending part of the loop of Henle. Clinical features of patients are compatible not only with PBS but also with true Bartter syndrome due to poor response to therapy and failure to a reduced capillary density in 9 patients out of 13 (Table 1).

In published maps and institutional affiliations.

### Table 1 (abstract A55). Results

| Capillaroscopic changes | N (15) | % |
|-------------------------|--------|---|
| Pathological capillaroscopy | 13 | 86.6 |
| Reduced capillary density | 5 | 38.4 |
| Tortuosity of the loops between 20-50% | 9 | 69.2 |
| Dilatation of the sub papillary plexus | 5 | 38.4 |
| Moderate angiotectonic disorder | 10 | 76.9 |

### A55
**Cystic Fibrosis and alterations of the peripheral microcirculation**

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### A56
**Functional endoscopic sinus surgery and pathogens detection in cystic fibrosis airways**

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### Background

Airways colonization with multiresistant pathogens such as P. aeruginosa (PA), S. maltophilia (SM), S. marcescens (SEM) and Methicillin-Resistant Staphylococcus aureus (MRSA) in cystic fibrosis increases the frequency of exacerbations, accelerate lung function decline and affect prognosis. Treating such pathogens is a challenge in cystic fibrosis management and nasal polyposis may work as bacterial reservoir. The aim of our study was to investigate the effect of functional endoscopic sinus surgery (FESS) on pathogens detection in cystic fibrosis airways.

### Materials and methods

In a retrospective observational study, we selected all subjects with diagnosis of cystic fibrosis with nasal polyposis who underwent a FESS in our center in Rovereto hospital. We cultured samples from lower airways (sputum) before and after surgery and samples from paranasal sinuses at surgery date. We compared: 1. the detection of multiresistant bacteria (PA, SM, SEM and MRSA) in lower airways and in paranasal sinuses, 2. the multiresistant pathogens in sputum before and after surgery.

### Results

The study included 11 subjects. We detected PA in 8 subjects, SM in 4 subjects and SEM in 2 sunjects, no MRSA was detected. In the first analysis we observed different pathogens in lower airways and in sinuses samples in 8 out of 11 subjects: we detected SM (4 subjects) and SEM (2 subjects) in samples from lower airways only and in other 2 subjects in paranasal sinuses only. In the second analysis we observed that in 3 subjects we could not detect multiresistant pathogens before surgery, two of them became positive after surgery.8 subjects were positive before surgery, 4 of them became negative after FESS. The paired t test showed a decrease of 27% (95CI -71/+16 %, p-value 0.17) of probability in detection of multiresistant bacteria after FESS.

### Conclusions

We observed a discrepancy in bacterial detection using samples from lower airways and paranasal sinuses. Moreover, our study shows a negative trend in detection of multiresistant bacteria in sputum after surgery. This might suggest a positive effect of FESS in controlling pathogen colonization in cystic fibrosis.

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