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To cite this version:

Sabine Bosselaar, Lucile Dhelin, Ellena Dautel, Marie Titecat, Stéphanie Duthoy, et al.. Taxonomic and phenotypic analysis of bifidobacteria isolated from IBD patients as potential probiotic strains.. BMC Microbiology, 2024, BMC Microbiology, 24, pp.233. 10.1186/s12866-024-03368-4. hal-04675360

HAL Id: hal-04675360 <https://hal.univ-lille.fr/hal-04675360v1>

Submitted on 22 Aug 2024

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Taxonomic and phenotypic analysis of bifdobacteria isolated from IBD patients as potential probiotic strains

Sabine Bosselaar^{1,2*}, Lucile Dhelin¹, Ellena Dautel¹, Marie Titecat², Stéphanie Duthoy¹, Marie Stelmaszczyk¹, Nathan Delory¹, Madeleine De Sousa Violante¹, François Machuron¹, Hassina Ait-Abderrahim¹, Pierre Desreumaux^{2,3}, Benoit Foligné² and Céline Monnet¹

Abstract

Background Infammatory Bowel Diseases (IBD) are a major public health issue with unclear aetiology. Changes in the composition and functionality of the intestinal microbiota are associated with these pathologies, including the depletion of strict anaerobes such as *Feacalibacterium prausnitzii*. Less evidence is observed for depletion in other anaerobes, among which bifdobacteria. This study characterized the taxonomic and functional diversity of bifdobacteria isolated from the human intestinal microbiota in active and non-active IBD patients by a culturomics approach and evaluated if these bifdobacteria might be used as probiotics for gut health.

Results A total of 341 bifdobacteria were isolated from the intestinal microbiota of IBD patients (52 Crohn's disease and 26 ulcerative colitis patients), with a high proportion of *Bifidobacterium dentium* strains (28% of isolated bifidobacteria). In ulcerative colitis, the major species identifed was *B. dentium* (39% of isolated bifdobacteria), in active and non-active ulcerative colitis. In Crohn's disease, *B. adolescentis* was the major species isolated from non-active patients (40%), while similar amounts of *B. dentium* and *B. adolescentis* were found in active Crohn's disease patients. The relative abundance of *B. dentium* was increased with age, both in Crohn's disease and ulcerative colitis and active and non-active IBD patients. Antibacterial capacities of bifdobacteria isolated from non-active ulcerative colitis against *Escherichia coli* LF82 and *Salmonella enterica* ATCC 14028 were observed more often compared to strains isolated from active ulcerative colitis. Finally, *B. longum* were retained as strains with the highest probiotic potential as they were the major strains presenting exopolysaccharide synthesis, antibacterial activity, and anti-infammatory capacities. Antimicrobial activity and EPS synthesis were further correlated to the presence of antimicrobial and EPS gene clusters by in silico analysis.

Conclusions Diferent bifdobacterial taxonomic profles were identifed in the microbiota of IBD patients. The most abundant species were *B. dentium*, mainly associated to the microbiota of ulcerative colitis patients and *B. adolescentis*, in the intestinal microbiota of Crohn's disease patients. Additionally, the relative abundance of *B. dentium* signifcantly increased with age. Furthermore, this study evidenced that bifdobacteria with probiotic potential (antipathogenic activity, exopolysaccharide production and anti-infammatory activity), especially *B. longum* strains, can be isolated from the intestinal microbiota of both active and non-active Crohn's disease and ulcerative colitis patients.

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Background

Infammatory Bowel Diseases (IBD), including Crohn's disease (CD) and ulcerative colitis (UC) are multifactorial disabling diseases representing a major public health issue. Both diseases represent a chronic relapsing–remitting condition afecting the gut. Scoring systems based on clinical, biological, endoscopic, radiologic and histologic observations have been developed and validated to describe disease activity; however, the defnitions of relapse and remission remain still under debate [1]. IBD have unclear aetiology implicating genetic susceptibility, environmental factors, and infuence of gut microbiota, contributing to a chronic infammation of the gastrointestinal tract $[2-4]$. Hence, the intestinal microbiota has gained much attention as an important factor in disease development, but also as a focus for new emerging therapeutic approaches with the development of probiotics, prebiotics or synbiotics.

Dysbiosis observed in the intestinal microbiota of IBD patients may play an important role in IBD pathogenesis. The intestinal microbiota of IBD patients have shown decreases in *Firmicutes* such as *Faecalibacterium prausnitzii*, *Blautia faecis* and *Roseburia inulinivorans*, increases in *Proteobacteria*, mainly enterobacteria such as *Escherichia coli* (with an increase in pathobiontic Adherent Invasive *E. coli*), and also *Fusobacterium* spp*., Ruminococcus gnavus* or pathogens such as *Clostridium difficile, Campylobacter* or *Salmonella*. This dysbiosis leads to functional diferences as bacteria with benefcial activities such as short chain fatty acid production and anti-infammatory capacities are decreased while bacteria with opportunistic detrimental functions such as pro-infammatory capacities or mucolytic activity are increased [5–7].

Generally, the intestinal microbiota of IBD patients are enriched in aerobic or aerotolerant bacteria and are depleted in strict anaerobes and it has been largely shown that *F. prausnitzii* is reduced in the intestinal microbiota of patients with IBD[5, 6]. Nevertheless, less evidence is made for other benefcial anaerobic bacteria, such as bifdobacteria. Several studies have shown decreased levels in bifdobacteria in IBD [8], while other increased abundance of bifdobacteria in UC (but not CD) compared to healthy controls [9–11]. To our knowledge, only very limited data was published focusing specifcally on bifdobacteria analysis within the microbiota of IBD patients at species level. Gueimonde et al*.* characterized intestinal adherent strains in colon cancer, IBD and diverticulitis patients and detected bifdobacteria in all IBD samples with a majority of *B. longum* and *B. bifdum* [12]. Furthermore, Duranti et al*.* identifed a reduction in *B. bifdum* in the intestinal microbiota of ulcerative colitis [13]. More studies investigated the probiotic efects of bifdobacteria in IBD. In diferent animal colitis models, several bifdobacteria strains reduced symptoms, improved histological colon observations and reduced the pro-inflammatory response (reviewed in $[14, 15]$). Human clinical trials also evidenced beneficial roles of supplementation of *B. longum* and *B. breve* strains in UC [16-19] and CD [20], with decreased disease activity, decreased endoscopic infammation and anti-infammatory efects. While bifdobacteria are known as one of the major known benefcial bacteria in the human intestinal microbiota and are largely studied for their probiotic application in IBD, the natural occurrence of bifdobacteria in IBD microbiota has not been fully characterized.

The aim of this in-depth bifidobacteria characterization in IBD patients by a culturomics approach was twofold: evaluate the taxonomic and functional diversity of bifdobacteria in active and non-active IBD patients and reveal if these bifdobacteria could potentially be used as probiotics in gut health. This work presents a culturomics approach to characterize the bifdobacteria microbiota of IBD patients in faecal samples collected during medical care at Lille University Hospital. Most studies profling dysbiosis in IBD were conducted by metagenomic approaches. These techniques give a good overall insight into microbiota composition but lack specificity for characterization of minor genera at species level and present many biases depending on method of analysis [21, 22]. Here, a culturomics approach allows us to focus on the bifdobacterial subpopulation of the intestinal microbiota, to characterize their functional potential in vitro and identify potential probiotic strains. We focused on three benefcial health efects known in bifdobacteria for functional characterization: exopolysaccharide (EPS) production, anti-pathogenic capacities, and immunomodulatory efects. Indeed, bifdobacteria produce a wide range of exopolysaccharides, mainly composed of galactose, glucose and rhamnose, with shown benefcial health efects such as protective efects favouring persistence in the gut, modulation of the microbiota, anti-pathogenic activity and immunomodulatory effects [23, 24]. We evaluated anti-pathogenic activity of bifdobacteria isolated from IBD patients on a IBD associated pathobiont, *E. coli* LF82 [25–27], but also on one of the major foodborne pathogens, *Salmonella enterica* [28, 29]. Anti-infammatory potential was evaluated by measuring decreased synthesis of IL-8 (a pro-infammatory chemokine activating neutrophils) in intestinal epithelial HT-29 cells, an extensively

recognised screening model for anti-infammatory probiotics [30–33]. A reduction in IL-8 synthesis is of interest in IBD patients as IL-8 increase has been shown in active UC and CD [34, 35] but also in global gut health, as IL-8 secretion is also increased in enteropathogeninduced infammation [36, 37]. We also established bacterial ftness and oxygen tolerance to identify if bacteria isolated from intestinal infamed environments could have adapted to their environment and present high tolerance levels and ftness and thus be particularly suitable as future probiotic candidates regarding industrial processes.

Methods

IBD patients

This study included patients with medical follow-up by a gastroenterologist at the Lille University Hospital (France) between the 2nd February 2021 and the 28th June 2022. Patients diagnosed with CD or UC, without antibiotic treatment in the last month, that provided faeces for medical care were included in this study. Faeces were collected for research purpose under secondary use with informed consent of the patients. For all samples, a database containing the following metadata was created: Pathology, age, sex, duration of pathology, location of infammation, symptomatic observations, and fecal calprotectin quantifcation. Patients were considered in active CD or UC if fecal calprotectin quantification was \geq 250 µg/g (with or without clinical symptoms) and in non-active CD or UC when fecal calprotectin quantifcation was < 250 µg/g and patients did not present symptomatic clinical gastrointestinal disorders. One patient with low fecal calprotectin quantification $\left(< 250 \text{ µg/g} \right)$ but presenting gastrointestinal disorders was excluded from analysis.

Analysis was performed on a total of 138 samples collected from 103 IBD patients: 98 samples were collected from CD (66% women, age 43.5 ± 13.6) and 40 from UC (65% women, age 49.6 ± 16.7), from patients with non-active $(CD = 59, UC = 22)$ and active disease $(CD = 39, UC = 18)$. Diversity of patients per pathology is described in Table 1.

Bacteria enumeration and strain isolation of fecal samples

Fecal samples were treated within 5h after collection. One g of fecal material was homogenized in 9mL of freshly regenerated cysteine diluent (heated at 90°C for 20 min and cooled down right before use). Serial dilutions were plated on freshly prepared Columbia Beerens medium [38], for bifdobacteria enriched enumeration and Columbia Blood Agar for total anaerobic bacteria enumeration*.* Plates were incubated at 37°C for 6 days in anaerobic jars. From Columbia Beerens agar plates, up to 10 colonies were randomly selected per fecal sample, on plates presenting isolated colonies. When diferent morphologies were picked, number of colonies picked per morphology were proportional to the representativity of this morphology in the sample to minimize biases in diversity ratios. Strains were then subcultured 3 times on agar plates from an isolated colony to guarantee purity on De Man-Ragosa-Sharpe (MRS) medium (Biomérieux, France) supplemented with 0.05% L-cysteine HCl at pH 6.5 (MRSc) in anaerobic conditions (Don Withley Anaerobic Workstation, 37°C, 10% H₂, 10% CO₂, 80% N₂).

Strain identifcation

Strains were taxonomically identifed by Matrix Assisted Laser Desorption Ionization—Time of Flight (MALDI-TOF) [39]. Bacterial isolates were cultivated on MRSc agar plates. Protein extraction procedure was performed on each isolate. $1 \mu L$ of the extract were deposited on a 96-well target plate and analyzed with the MALDI-TOF mass spectrometer according to the manufacturer's recommendations (Brüker, Wissembourg, France). Species were identifed using an accurate identifcation $score > 1.9$.

Identifcation was confrmed by Whole Genome Sequencing. Total genomic DNA was extracted from pure bacterial cultures using EZ-96 isolation kit tissue DNA from Omega Biotek following provider instructions with exception of the lysis step. Lysis was achieved by enzymatic reaction (20 mM Tris, 2 mM EDTA, pH $8+20$ mg/mL lysozyme+1,2% Triton, incubation 30 min at 30 °C) prior to starting the provider process. Sequencing libraries were prepared using Illumina DNA Prep kit following providers instructions. Isolated DNA and libraries samples were quantifed and validated using the Qubit fuorometric method (Invitrogen). Sequencing of an equimolar pool of multiplexed samples was realized on a Nextseq 550 lllumina sequencer in paired-end sequencing mode producing reads of 149 bases using PhiX as control. Illumina short reads were de novo assembled using Spades (v3.15), and quality controls were performed on all genomes with Quast (v5.1). Species were identifed using BLAST on the whole genome GTDB database and assessed using the ANI (Average Nucleotide Identity) metric calculated by fastANI (>96% to type strain) [40]. To check species identifcation, a phylogenetic tree was created by unweighted paired group mean arithmetic (upgma) method using MASH distances and represented using ggtree.

Crohn's Disease Ulcerative Colitis Non active Active All CD patients Non active Active All UC patients Number of samples 59 (60%) 39 (40%) 98 (100%) 22 (55%) 18 (45%) 40 (100%) Sex Men 22 (37%) 11 (28%) 33 (34%) 10 (45%) 4 (22%) 14 (35%) Women 37 (63%) 28 (72%) 65 (66%) 12 (55%) 14 (78%) 26 (65%) **Age** <50 years 38 (64%) 20 (51%) 58 (59%) 10 (45%) 13 (72%) 23 (58%) ≥50 years 21 (36%) 19 (49%) 40 (41%) 12 (55%) 5 (28%) 17 (42%) **Gastro-intestinal disorders** Yes 16 (41%) 16 (16%) 9 (50%) 9 (23%) No 59 (100%) 22 (56%) 81 (83%) 22 (100%) 8 (44%) 30 (75%) Unknown 1 (3%) 1 (1%) 1 (6%) 1 (2%) **Location of infammation** Small intestine only 19 (68%) 19 (49%) 59 (60%) Colon 19 (32%) 20 (51%) 39 (40%) 22 (100%) 18 (100%) 40 (100%) **Duration of pathology** <5 years 6 (10%) 7 (18%) 13 (13%) 2 (9%) 2 (11%) 4 (10%) 5–15 years 27 (46%) 11 (28%) 38 (39%) 11 (50%) 10 (56%) 21 (53%) 15–25 years 14 (24%) 12 (31%) 26 (27%) 5 (23%) 3 (17%) 8 (20%) ≥25 years 11 (19%) 3 (8%) 14 (14%) 4 (18%) 2 (11%) 6 (15%) Unknown 1 (2%) 6 (15%) 7 (7%) 1 (6%) 1 (2%) **Strain isolation** Samples with bifdobacteria isolated 29 (49%) 23 (59%) 52 (53%) 14 (64%) 12 (67%) 26 (65%) Samples without bifidobacteria isolated 30 (51%) 16 (41%) 46 (47%) 8 (36%) 6 (33%) 14 (35%) **Total number of strains isolated 379 (63%) 219 (37%) 598 (100%) 122 (50%) 121 (50%) 243 (100%) Strain isolation** Total number of bifdobacteria isolated 131 (35%) 104 (47%) 235 (39%) 48 (39%) 58 (48%) 106 (44%)

Table 1 Characteristics of sample origin (*N*=138 fecal samples) from IBD patients

In vitro functional characterization

For in vitro functional characterization, bifdobacteria were routinely grown in liquid MRSc, autoclaved and stored under N_2 atmosphere. Bifidobacteria were inoculated at 10% in a 96-well DeepWell, precultured for 24h, Optical Density at 620nm (OD_{600nm}) adjusted at 0.08 and cultured for another 18h to reach the end of exponential growth phase. Bifdobacteria were routinely incubated at 37°C in anaerobic conditions (10% H_2 , 10% CO₂, 80% N₂, Bact-R Plus System).

Growth characterization and oxygen tolerance

To determine growth curves, OD_{600nm} adjusted precultures were incubated in anaerobic conditions (10% H_2 , 10% $CO₂$, 80% N₂, Don Withley Anaerobic Workstation) and OD_{620nm} was measured every 20min (Byonoy Absorbance 96 plate reader). Growth parameters were calculated with the Growthcurver R package [41].

To determine oxygen tolerance, OD_{600nm} adjusted precultures were incubated in microaerophilic conditions (5% of residual O_2 , Bact-R Plus Systems) and OD_{600nm} was read after 24h and 48h of incubation at 37°C. To evaluate oxygen tolerance, ratio of OD_{600nm} in microaerophilic conditions / OD_{620nm} in anaerobic conditions was calculated for each strain. Strains with ratios>0.6 were considered tolerant to growth in microaerophilic conditions and strains with ratios<0.2 were considered intolerant to growth in microaerophilic conditions.

Exopolysaccharides production

EPS producing strains were identifed by their mucoid phenotype when grown on agar plates as described by Ruas Madiedo [42]. Briefy, bifdobacteria precultures were spotted on 0.05% L-cysteine HCl—MRS agar plates with 20g/L glucose, 60g/L glucose, 100g/L sucrose, 100g/L maltose or 100g/L lactose. Strains presenting a mucoid phenotype after 72h incubation (37 $^{\circ}$ C, 10% H₂, 10% $CO₂$, 80% N₂, Bact-R Plus System) were considered EPS producing strains. Strains forming white dry colonies were considered non-EPS producing strains. EPS

phenotype of strains presenting an intermediate phenotype ("classic" white colonies) were not established.

In silico, exopolysaccharide gene clusters were predicted for strains presenting an EPS producing phenotype in vitro. Priming glycosyltransferase encoding genes were searched by BLAST against the B624_0342 gene from *B. longum* 35624 (accession number CP013673). Extended genomic regions around the priming glycosyltransferase gene were manually analyzed to identify potential EPS gene cluster regions. Genetic regions were annotated by combining genomic annotations from Prokka [43]. BLAST against amino acid sequences of EPS related genes described in other bifdobacterial gene clusters [23, 44, 45] and InterPro [46].

Antibacterial activity

To test the antibacterial activity, co-culture spot plate assays were set up, similarly to described by O'Riordan and Fitzgerald [47]. *Bifdobacterium* precultures were spotted on MRS (Biomérieux, France) agar plates supplemented with 0.05% L-cysteine HCl pH 6.5 and grown for 24h at 37°C in anaerobic conditions (37°C, 10% H₂, 10% $CO₂$, 80% N₂, Bact-R Plus System) before addition of an overlay of *Salmonella enterica subsp. enteriditis* ATCC 14028 (inoculation at OD_{600nm} 0.005) or *E. coli* LF82 (inoculation at \rm{OD}_{600nm} 0.05) in soft agar (8g/L agar) TSB or LB medium respectively. After a supplementary 24h incubation (37°C, 37°C, 10% H₂, 10% CO₂, 80% N₂, Bact-R Plus System), inhibition scores were attributed according to size of inhibition zones (from 0.25 for nearly visible inhibition zones to 1 for control strain *B. longum* DSM 20219). Mean inhibition scores of replicates were calculated, and inhibition was considered strong for average scores above 0.65 and absent for scores below 0.35.

In silico, antimicrobial gene clusters were predicted using BAGEL4 (v4.0) for bacteriocins and ribosomally synthesized and post-translationally modifed peptides and antiSMASH (v6) for other secondary metabolite [48, 49].

Immunomodulatory efect

Immunomodulatory effect was evaluated on a subset of 130 strains. For this selection, groups of genetically identical strains (fastANI>99.99%) were formed and one representative per group was included. Immunomodulatory efect was evaluated by measuring reduction of LPS-stimulated IL-8 secretion by bifdobacteria strains in a HT-29 cell culture model. This model was largely described in literature [33, 50–58] with diferent stimulants at various concentrations to induce infammatory response and diferent incubation times. Here, the model was adapted for high throughput screening to a 96-well MTP assay and optimal amount of HT29 cells per well, cell/bacteria ratio, LPS concentration and incubation times were defned after multiple tests during method set-up. HT-29 cells (HTB-38, ATCC) were routinely cultivated in DMEM 4,5g/L D-glucose, pyruvate, GlutaMAX, 10% FCS (Gibco) at 37°C, 5% CO₂. Bifidobacteria cultures were adjusted to $OD_{600nm} = 0.25$ (approximately 4.10^8 CFU/mL) and diluted 20-fold in DMEM medium. Bacterial cell suspensions were added on previously plated and adhered HT-29 cells (40 000 cells / well in 96 well plates) at MOI 50 and incubated for 4h (37°C, 5% CO₂). After 4h incubation, inflammation was induced by addition of 10ng/mL of lipopolysaccharide (LPS) 0111:B4 (L4391, Sigma-Aldrich). Supernatants were collected after 24h incubation (37°C, 5% CO2), flter sterilized (0.2µm) and stored at -80°C until analysis. IL-8 secretion by HT-29 cells was measured by ELISA (Duo-Set ELISA Development System Human IL-8/CXCL8, DY208—R&D Systems) according to the manufacturer's protocol. For inter-experiment normalization, results were expressed as diminution of IL-8 secretion by bifdobacteria compared to control condition (LPS-stimulated HT-29 cells without probiotic treatment).

Statistical analysis

The number of isolated strains summarized by families were presented using percentages to consider the unbalanced numbers of patients in groups. Percentages were compared between groups using $Chi²$ tests. In case of overall significant *p*-value, post-hoc Chi² tests were performed for each family with adjustment of *p*-values with Bonferroni method to control type-1 error infation due to multiplicity testing. For functional characterization, all experiments were performed in technical and biological duplicates and mean results on the 4 values were considered for analysis.

Results

Bifdobacteria isolation in IBD patients *Similar fecal sample enumerations between active and non‑active CD and UC*

Bacteria of fecal samples were enumerated on Columbia Beerens (*Bifdobacteriaceae* enrichment) and Columbia blood agar (Total anaerobic flora). No significant differences were observed in bacterial enumerations between samples from patients in CD and UC (Wilcoxon test *p*-value=0.38 for Columbia Beerens Agar and 0.43 for Columbia Blood Agar). In CD and UC patient groups, no signifcant diferences were observed in bacterial enumerations (*Bifdobacteriaceae* enrichment or total bacteria) between active or non-active patients (Wilcoxon test *p*-values > 0.05, Fig. 1).

Fig. 1 Bacterial enumeration of fecal samples on Columbia Beerens medium (Bifdobacteriaceae enrichment) and Columbia blood agar (Total bacteria) grouped by pathology and disease activity. Boxplots represent median (solid line), frst and third quartile (lower and upper boundaries), lower and upper limits (whiskers, defned by 1.5* inter quartile range) and outliers (dotpoints). Wilcoxon test *p*-values: *: 0.01–0.05; **: 0.001–0.01; ***:<0.001; ns:>0.05. N: number of samples per group

Bifdobacteria isolated from IBD were enriched in *B. dentium* **and** *B. adolescentis*

A total of 841 strains (from 136 fecal samples) were isolated on Columbia Beerens medium, among which 341 were identifed as bifdobacteria. For 58 fecal samples, only lactic acid bacteria were isolated. From the other 78 fecal samples (52 CD and 26 UC), 341 bifdobacteria were isolated. Bifdobacteria were isolated from UC and CD patients in similar ratios, with respectively 39 and 44% of total isolated strains in these patient groups identifed as bifdobacteria. Fecal microbiota of IBD patients were enriched in *B. dentium* (27.3% of all bifdobacteria isolated, *n*=93 strains), *B. adolescentis* (25.5%, *n*=87 strains) and *B. longum* (17%, *n*=58 strains). Other species isolated were *B. bifdum* (14%), *B. pseudocatenulatum* (12%), *B. breve* (2%), *B. angulatum* (2%) and *B. catenulatum* (1%) (Fig. 2).

Bifdobacteria profles difer between CD and UC and vary between remission and relapse

Bifdobacteria profles difered between CD and UC patients (Chi² test *p*-value<0.001) due to different ratios of *B. adolescentis* (adjusted *p*-value<0.001) and *B. dentium* (adjusted *p*-value<0.05). In CD, a majority of *B. adolescentis* strains were isolated (33%) while in UC, a majority of *B. dentium* were isolated (39%) (Fig. 3A).

We further analyzed if these diferences in bifdobacteria profles could be due to the location of the infammation in the digestive tract. No signifcant diferences were identifed in bifdobacteria profles between CD patients where infammation was located only in the small intestine or also afected the colon (Chi² test *p*-value=0.1, adjusted *p*-values per species=1). Diferent bifdobacteria profles were observed between UC patients and colic forms of CD (Chi² test *p*-value < 0.001) with increased *B. adolescentis* ratios in UC (adjusted *p*-value < 0.001). In UC, most strains isolated were *B. dentium* (39%) and only 10% were *B. adolescentis*. In contrast, in colic forms of CD, the majority of strains isolated were identifed as *B. adolescentis* strains (34%) and only 24% as *B. dentium* (data not shown). These observations were identical to global observations between UC and CD (Fig. 3A).

Bifdobacteria taxonomic profles difered between active and non-active CD (Chi² test p -value=0.014) but not active and non-active UC (Chi² test p -value > 0.05) (Fig. 3B). Nevertheless, adjusted *p*-values did not highlight any signifcant diferences for specifc bifdobacterial species between non-active and active CD. The same bifidobacterial taxonomic profles were identifed in active CD and UC (Chi² test p -value > 0.05) while non-active CD and UC significantly differed (Chi² test p -value < 0.001) in *B. adolescentis* (adjusted *p*-value<0.001) and *B. dentium* (adjusted *p*-value<0.01). Indeed, *B. adolescentis* was predominant in non-active CD (40% of bifdobacteria isolated), whereas *B. dentium* was predominant in active CD (28%), and active (34%) and non-active UC (44%) (Fig. 3B).

Fig. 2 Taxonomic species identifcation of the 341 bifdobacteria strains isolated from 78 IBD patients (*N*=52 Crohn's disease patients+*N*=26 Ulcerative colitis patients). Ratios are presented as relative abundance of bifdobacteria isolated (absolute number of bifdobacteria strains isolated)

Fig. 3 Characterization of the bifdobacteria profle in intestinal microbiota of IBD patients by (**A**) pathology or (**B**) disease activity. Ratios are presented as relative proportion of bifdobacteria isolated (absolute number of strains isolated). Ratios were not displayed for species with relative abundance ≤ 5%. Chi² test *p*-values: *: 0.01-0.05; ***: 0.001-0.01; ***: < 0.001; ns: > 0.05. N: number of samples from which strains were isolated. n: number of strains isolated

Fig. 4 Characterization of the bifdobacteria profle in intestinal microbiota of IBD patients by age (**A**) for all patients, (**B**) per pathology and (**C**) per disease activity. Ratios are presented as relative proportion of bifdobacteria isolated (absolute number of bifdobacteria strains isolated). Ratios were not displayed for species with relative abundance ≤5%. N: number of samples from which strains were isolated. n: number of strains isolated. Chi.² test *p*-values: *: 0.01–0.05; **: 0.001–0.01; ***:<0.001; ns:>0.05

Age-related evolution in bifdobacteria profles in IBD patients

In IBD patients, relative abundances of bifdobacteria species varied with age $(Chi^2$ test p -value<0.001 between patients < and \geq 50 years old), notably with a signifcant increase in *B. dentium* in IBD patients above 50 years old, and a signifcant decrease in *B. longum* and *B. pseudocatenulatum* (adjusted *p*-values<0.001 between groups<50 and \geq 50 years old) (Fig. 4A). Increase in *B*. *dentium* with age was observed in UC and CD (adjusted *p*-values<0.001, Fig. 4B) and in non-active (adjusted p -values < 0.01) and active (adjusted p -values < 0.001) IBD patients (Fig. 4C). In the same manner, decrease in *B. longum* with age was observed in UC (adjusted *p*-values < 0.001) and CD (adjusted p -values < 0.01, Fig. 4B) and in non-active (adjusted p -values < 0.01) and active (adjusted *p*-values < 0.001) IBD patients (Fig. 4C). These observations did not seem to correlate with duration of the pathology. Mean age of patients was not signifcantly diferent per pathology groups (Wilcoxon test p -value = 0,3) and differences in microbiota composition with age did not seem to correlate with microbiota variations according to the duration of the pathology (data not shown).

Functional characterization of bifdobacteria isolated from IBD patients

In vitro growth capacities

As bifdobacteria were isolated from intestinal infammatory environments, bacteria ftness was evaluated by determining anaerobic growth kinetics and oxygen tolerance to evidence possible diferences according to isolation origin. In vitro growth parameters were evaluated by following growth kinetics in anaerobic and microaerophilic $(5\%$ O₂) conditions. Growth abilities largely difered between strains, but trends were observed per species. Globally, in anaerobic conditions, *B. longum* presented the best growth abilities with reduced lag phases (average of 6.5h), highest growth rates (average of 0.36 h^{-1}) and highest OD_{600nm} reached (average of $OD_{600nm} = 0.9$). For other species, lag phases were around 10h, growth rated around 0.25h⁻¹ and maximum OD_{600nm} were heterogeneously distributed among strains, but generally reduced for *B. bifdum* strains $(OD_{600nm} = 0.4 \text{ on average})$ (Fig. 5A).

In microaerophilic conditions (5% O_2), 100% (*n*=58 strains) of *B. longum*, 100% (*n*=6) of *B. breve*, 88% (*n*=36) of *B. pseudocatenulatum* and 70% (*n*=33) of *B. bifdum* presented equivalent growth capacities to anaerobic conditions. *B. adolescentis* and *B. dentium* showed limited oxygen tolerance with respectively 55% (*n*=48) of *B. adolescentis* strains and 37% (*n*=34) of *B. dentium* strains exhibiting no growth in microaerophilic conditions (Fig. 5B).

Bacterial ftness in anaerobic conditions and oxygen tolerance did not depend on isolation origins: no signifcant diferences were found between strains isolated from CD or UC patients in active or non-active disease.

Exopolysaccharide synthesis

EPS secretion was evaluated by observing the aspects of colonies on agar plates containing diferent carbohydrate sources. EPS synthesis was strain dependent with a wide variety of phenotypes within the same species. None of the *B. bifdum* (*n*=47) and only one *B. dentium* (*n*=93) tested presented an EPS producing phenotype on the 20g/L glucose MRS agar plates. *B. longum* strains had high EPS producing potential, with 24% (*n*=14) of

Fig. 5 Growth characterization of bifidobacteria in (A) anaerobic and (B) microaerophilic (under 5% 0₂) conditions. A Growth parameters were not determined for strains with Max OD_{600nm} < 0.2 or that did not reach end of exponential phase after 48h. Boxplots represent median (solid line), first and third quartile (lower and upper boundaries), lower and upper limits (whiskers, defined by 1.5* inter quartile range) and outliers (dotpoints). (B) Not determined: strains that did not present growth in control condition (anaerobic) to calculate microaerophilic growth ratio

Fig. 6 EPS production on 20g/L glucose MRS agar plates (**A**) per bifdobacterial species and (**B**) according to strain origin (pathology and disease activity). Chi.² test *p*-values: *: 0.01–0.05; **: 0.001–0.01; ***: < 0.001; ns: > 0.05

B. longum strains producing EPS. We also identifed 9 *B. adolescentis*, 1 *B. angulatum*, 1 *B. breve*, 1 *B. catenulatum* and 7 *B. pseudocatenulatum* EPS producing strains

(Fig. 6A). Similar results were obtained for EPS synthesis when growing on other carbohydrate sources i.e. lactose, maltose and sucrose (data not shown).

For 32 out of the 34 strains with an EPS producing phenotype, potential EPS encoding gene clusters were detected in silico. All genomic clusters contained a priming glycosyltransferase gene: *rfbP*, encoding an undecaprenyl-phosphate sugar phospho-transferase were mainly identifed in *B. adolescentis*, *B. pseudocatenulatum* and *B. dentium* while *cpsD*, a galactosyl-transferase, was predominant in *B. longum*, *B. angulatum* and *B. catenulatum* (Supplementary Figure S1). The genetic regions surrounding the primary glycosyltransferase gene displayed diferent structural organizations, but contained characteristic genes involved in EPS synthesis (including glycosyltransferases, genes involved in rhamnose biosynthesis and transport related fippases or ABC transporters). In all 13 *B. longum* EPS gene clusters, several glycosyltransferases and chain length determinant proteins were identifed. Flippases or ABC transporter encoding genes were found in 5 EPS gene clusters and genes described as involved in polysaccharide biosynthesis in 8 EPS gene clusters. *B. pseudocatenulatum* strains also presented potential complete EPS gene clusters as the 7 genetic regions identifed all contained a priming glycosyltransferase, glycosyltransferases and other polysaccharide synthesis related genes. Five out of the 7 potential EPS gene clusters also presented a fippase encoding gene. In *B. adolescentis*, in the surrounding regions of a potential priming glycosyltransferase identifed in 9 strains, only one presented other glycosyltransferases and a polysaccharide biosynthesis related protein. For the other potential EPS encoding regions, a chain-length determinant protein like gene was identifed and in 6 regions, rhamnose biosynthesis genes were highlighted. Finally, potential complete EPS gene clusters were identifed in 1 *B. angulatum*, 1 *B. catenulatum* and 1 *B. dentium* strain, containing a priming glycosyltransferase, several glycosyltransferases, polysaccharide biosynthesis protein and transporter encoding genes (Fig. S1).

Regarding the origin of the strains, EPS production frequency in bifdobacteria did not correlate with isolation origin. No diferences were observed between strains isolated from active and non-active CD or from active and non-active UC (Chi2 test *p*-value>0.1) (Fig. 6B). Correlations were neither found according to other isolation origin parameters (age, length of pathology, overall differences between pathologies or disease activity).

Antibacterial activity

Antibacterial activity of the 341 bifdobacteria was evaluated by co-culture and inhibition halo observation for *Salmonella enterica* subsp. *enterica* ATCC 14028 and *Escherichia coli* LF82 strains (Fig. 7A, B). Most strains with high anti-pathogenic activity against both strains were *B. bifdum* (53% of strains inhibited both pathogens), *B. longum* (40%) and *B. dentium* (31%) while *B. adolescentis*, *B. angulatum*, *B. breve*, *B. catenulatum* and *B. pseudocatenulatum* strains had very limited antibacterial activity, with none or only 1 strain per species able to inhibit both pathogens (Fig. 7A, B).

Bifdobacterial strains inhibiting both *E. coli* and *S. enterica* were enriched in non-active UC compared to active UC (Chi² test p -value = 0.017). In UC, bifidobacteria inhibiting both *E. coli* (Chi² test p -value=0.025) and *S. enterica* (Chi² test *p*-value=0.037) were more prevalent in non-active disease compared to active disease. In CD, the same trend was observed but without statistical signifcance (Fig. 7 C, D). No correlations between antibacterial activity and other strain origin characteristics were identifed.

In addition to in vitro antimicrobial screening, biosynthetic gene clusters (BGCs) potentially involved in the synthesis of antimicrobial metabolites were searched by antiSMASH and BAGEL4 bioinformatic tools. BGCs were identifed in 115 genomes, corresponding to 34% of all strains (Fig. 8). Gene clusters identifed are represented in supplementary fgures S2 and S3. Mainly, a propionicin SM1 like gene cluster was identifed in 61 *B. dentium* genomes (66% of all *B. dentium* strains), 1 *B. longum* and 1 *B. adolescentis* genome. The lanthipeptide BLD_1648 gene cluster was detected in 8 *B. longum* and 2 *B. dentium* genomes. Other lanthipeptide gene clusters were identifed in 31 *B. dentium* (class I lanthipeptides) and 15 *B. pseudocatenulatum* (class IV lanthipeptides). Furthermore, pinensin-like and lactoccoccin like gene clusters were found respectively in 2 *B. longum* and 3 *B. pseudocatenulatum* genomes. Finally, 6 Non-Ribosomal Peptide Synthetase (NRPS) gene clusters were identifed in *B. adolescentis* genomes and 7 polyketide synthetases (PKS) in *B. breve* and *B. longum* strains.

Among the 25 bifdobacteria that presented strong inhibition against both *E. coli* and *S. enterica*, only 8 (6 *B. longum* and 2 *B. dentium*) presented BGCs. Among these strains, we fnd the 2 *B. longum* strains for which the pinensin 11 BGCs were identifed and the 3 *B. longum* with the hglE-KS BGCs. Furthermore, one *B. longum* strain with high antimicrobial activity encoded the BLD₋₁₆₄₈ gene cluster. The 2 *B. dentium* strains with high antimicrobial activity in vitro both encoded the propionicin SM1 gene cluster and one of them also presented an unknown class I lanthipeptide BGC. From the 56 strains inhibiting only one of the two pathogens in vitro, BGCs were identifed in 15 strains, among which 11 the propionicin SM1 and 4 the BLD_1648 gene cluster. In contrast, among the 155 strains that did not have antimicrobial activity in vitro against *E. coli* or *S. enterica*, 63 strains encoded for BGCs (Propionicin SM1, BLD_1648,

Fig. 7 Inhibition of E. coli LF82 and Salmonella enterica ATCC 14028 by bifdobacteria. **A** Inhibition profles by bifdobacteria per species and per pathogen tested. **B** Global inhibition profles by bifdobacteria species (number of pathogens inhibited). **C** Inhibition profles by bifdobacteria per pathogen tested according to isolation origin (pathology and disease activity). **D** Global inhibition profles according to isolation origin (pathology and disease activity). Chi² test *p*-values: *: 0.01–0.05; **: 0.001–0.01; ***: < 0.001; ns: > 0.05. n: number of strains. Not determined: insufficient growth of bifidobacterial strain in experiment conditions

Lactoccocin_972, diferent lanthipeptides, KPS and NRPS gene clusters). Further analysis will be needed to evaluate if these strains indeed produce these antimicrobial compounds in vitro, their exact structure and spectrum of action.

Immunomodulatory activity

Immunomodulation was determined by assessing inhibition of LPS-induced IL-8 secretion by HT-29 human epithelial cells after treatment with a selection of 105 genetically different bifidobacteria. Herein, from the seven bifidobacterium species isolated, only *B. longum* and *B. breve* strains presented significant anti-inflammatory activity. Only 4 strains (all *B. longum*) out of the 105 strains tested presented high anti-inflammatory activity (> 70% inhibition of IL-8 secretion compared to LPS-induced control). *B. longum* thus presented the highest anti-inflammatory activity (13% of *B. longum* strains tested presented high anti-inflammatory activity). An additional 45% of the *B. longum* (*n*=14) presented moderate anti-inflammatory activity (inhibition of 40 to 70% of IL-8 secretion) (Fig. $9A$).

Regarding the origin of the strains, no differences in anti-inflammatory activity were found between *B. longum* isolated from patients in active er non active disease (Chi² test *p*-value = 1, Fig. 9B) or between UC or CD (Chi² test *p*-value = 0.7, data not shown). Statistical analyses were not performed according to pathology and disease activity combined due to reduced strain number per group.

Fig. 8 Biosynthetic gene clusters identifed in bifdobacteria genomes by combined analysis with antiSMASH and BAGEL4

Discussion

To our knowledge, only few studies were published focusing specifcally on taxonomic diversity of bifdobacteria in intestinal microbiota of IBD patients at species level. These studies included a limited number of patients (<15 IBD patients), and do not distinguish active from non-active IBD [12, 13]. Here, we set up a culturomics approach to give insight into the taxonomic diversity of bifdobacteria in the IBD microbiota of a large group of patients (103 patients), in active (57 samples) or nonactive IBD (81 samples) (Fig. 10). As fecal samples were collected as secondary use as part of a medical followup, the study design included only IBD patients and no healthy controls, setting the focus of this work to comparisons between active and non-active CD and UC. We collected a large metadata set allowing us to analyze bifdobacteria microbiota in IBD patients according to diferent parameters such as type of pathology, disease activity, disease location, age or duration of pathology. We defned disease activity considering fecal calprotectin quantifcation (a specifc marker of gastrointestinal infammation) and symptomatic observations. Calprotectin levels are correlated with endoscopic activity but present limitations in terms of specifcity and individual variation. Fecal calprotectin levels described in the literature to predict disease activity, mucosal healing or disease relapse risk go from 50 μ g/g to 500 μ g/g, with sensitivities going from 61 to 100% and specifcities between 43 and 90% [59–61]. In our analysis, we defned non-active UC or CD as patients with fecal calprotectin levels strictly lower than $250 \mu g/g$ that did not experience any gastrointestinal disorders. Active CD or UC was defned as patients with fecal calprotectin levels equal to or higher than 250 µg/g (i.e. patients with infammation in the gastro-intestinal tract, with or without gastrointestinal disorders).

With this culturomics approach, bacterial enumeration on bifdobacteria enrichment medium and total anaerobic flora were similar between patients in CD and UC, regardless of disease activity. This was consistent with the fnding of Premkumar et al*.*, who did not identify differences in *Bifdobacterium* abundance between healthy

Fig. 9 Immunomodulatory activity of bifdobacteria by inhibition of IL8 secretion of HT29 cells. (**A**) Immunomodulatory profle of all bifdobacteria per species. (B) Immunomodulatory profile of B. longum according to disease activity. Chi² test *p*-values: *: 0.01–0.05; **: 0.001–0.01; ***: < 0.001; ns:>0.05. n: number of strains

subjects, CD and UC patients by qPCR [62]. Nevertheless, after taxonomic identifcation of isolated strains, only 41% of isolated strains appeared to be bifdobacteria and for 44% of the samples, no bifdobacteria were isolated. Isolates were randomly selected from Columbia Beerens agar plates, but as this medium was not fully selective for bifdobacteria, some minor bifdobacteria populations in fecal samples might not have been evidenced here, especially in the case of a high abundance of lactic acid bacteria with favored growth on Columbia Beerens Medium. This created a bias in our study due to the culturomics approach chosen, in which only part of all cultivable bifdobacteria present might have been identifed.

Diferences in bifdobacteria microbiota composition according to pathology and IBD activity in this study were observed mainly between the two major species isolated, *B. adolescentis* and *B. dentium*, 2 closely related species belonging to the *B. adolescentis* phylogenetic group [63]. *B. dentium* were enriched in UC, both in non-active and active disease. In contrast to our results, studies of Gueimonde et al. and Duranti et al., analyzing bifdobacteria profles in IBD patients, did not identify *B. dentium* strains [12, 13]. We also highlighted increased *B. dentium* relative abundances in patients above 50 years old. These results are consistent with studies on healthy subjects showing increased *B. dentium* in elderly [64, 65]. *B. dentium* are known as members of the human intestinal microbiota but are not generally found as the major bifdobacteria species present. Furthermore, *B. dentium* was frst identifed in dental caries and several studies confrmed their cariogenic properties (ability to survive in the human oral cavity and acidogenic potential) [66, 67]. Associations were established between IBD and oral diseases (periodontitis, higher caries incidence) [68–70], which could correlate with the high *B. dentium* ratios found in IBD patients in our study. Nevertheless, *B. dentium* are also part of the human healthy microbiota and have shown benefcial efects in animal models of IBD. In vitro, *B. dentium* strains have been shown to positively infuence the epithelial barrier function by decreasing gut permeability $[71–74]$, and regulating mucin synthesis [75]. In vivo, strains of *B. dentium* were able to relieve colitis symptoms in a DSS stimulated mice model [72], decreased weight loss and levels of TNF, IL-8 and IL-6 infammatory markers in a TNBS-induced colitis mice model [76]. These data suggest that higher abundance of *B. dentium* in IBD patients should not only be considered as a marker of dysbiosis but could play a benefcial role.

Furthermore, this study found increased *B. adolescentis* relative abundances in CD compared to UC, especially in non-active CD. This observation was not consistent with studies of Gueimonde et al. and Duranti et al., who did not reveal *B. adolescentis* as specifc markers of IBD [12, 13] while Barberio et al*.* showed increased *B. adolescentis* in non-active UC compared to active UC and healthy controls [77]. *B. adolescentis* is generally known as one of the main bifdobacterial species of the human

Fig. 10 Summary of bifdobacteria characterization in IBD patients

gut microbiota and largely studied for its potential probiotic applications. One of the most evidenced functionalities of *B. adolescentis* is γ-aminobutyric acid secretion, a major inhibitory neurotransmitter that could be responsible for anxiolytic and antidepressant efects. Furthermore, *B. adolescentis* were shown to secrete vitamin B9 and have benefcial efects on the gut barrier in in vitro colitis models by increasing mucus layer and present immunoregulatory efects (stimulation of regulatory T cells, reduction of nuclear factor κB) (reviewed in [78]). These beneficial characteristics of *B. adolescentis* might explain their prevalence in non-active CD.

The higher *B. dentium* and *B. adolescentis* abundances identifed in our analysis could be specifc to our population, with environmental and geographic infuences, or depend on the isolation method used. All bifdobacteria species presented sufficient growth on the selective medium used (Columbia Beerens medium) but no in-depth analysis was performed on the selectivity of this medium. However, our culturomics approach did not seem to favor oxygen tolerant species,*B. dentium* and *B. adolescentis*, as these species are among the most oxygen sensitive bifdobacteria in our in vitro functional characterization. Comparison of diferent studies

describing the taxonomic diversity of bifdobacteria in IBD remains challenging as other studies are mainly based on metagenomic approaches andalso include many biases during sample treatment and bioinformatic analysis $[21, 22]$. This complicates identification and semi-quantitative analysis for minor populations of the intestinal microbiota.

During functional characterization of our bifdobacteria collection, we confrmed previously known phenotypes and functions of bifdobacteria (oxygen tolerance, exopolysaccharide synthesis, antimicrobial activity, and immunomodulatory effect) on a large number of strains.

Exopolysaccharide production in vitro was associated with potential EPS gene clusters in diferent *B. longum*, *B. pseudocatenulatum* and *B. adolescentis* strains. *B. longum* and *B. pseudocatenulatum* presented EPS gene clusters similar to those already described in the literature [23, 44, 45], containing genes encoding for priming glycosyltransferases, glycosyltransferases, rhamnose biosynthesis, chain length determinant proteins and other polysaccharide biosynthesis proteins. Nevertheless, no glycosyltransferase encoding genes were identifed in *B. adolescentis* potential EPS gene clusters. Further in vitro and in silico research will be needed to correlate EPS producing phenotypes to gene clusters and identify if the genetic regions identifed here are related to EPS synthesis in *B. adolescentis* strains. Besides highlighting the antimicrobial activity of *B. longum* and *B. bifdum*, already described in literature, we evidenced probiotic potential of *B. dentium* strains through inhibition of *E. coli* LF82 and *S. enterica ATCC* 14028. Antimicrobial activity of the bifdobacterial strains can be associated with several mechanisms such as the production of acetic and lactic acids, but also synthesis of specifc antimicrobial compounds. 115 bifdobacterial genomes presented BGCs encoding potential bacteriocins, ribosomally synthesized and post-translationally modifed peptides or non ribosomally synthesized peptides (NRPS, PKS). Most noticeable, 66% of the *B. dentium* strains could potentially produce a propionicin SM1 like bacteriocin.

Most of the probiotic characteristics assessed were already largely described in literature, but herein, this study evaluated if these characteristics were associated with the isolates' origin (type of patient, disease activity, age, pathology duration). No diferences were observed in bacterial ftness, oxygen tolerance, EPS synthesis and immunomodulatory activity between isolation origin. Bifdobacteria isolated from non-active UC presented increased anti-pathogenic activity against both Adherent Invasive *E. coli* LF82 and *S. enterica* ATCC 14028 compared to bifdobacteria isolated from active UC. More generally, our study highlighted that bifdobacteria, even when isolated from infamed gastrointestinal environments from active IBD patients present potential probiotic characteristics. *B. longum* presented the highest ratios of strains for EPS synthesis, antibacterial activity, and anti-infammatory capacities. The potential probiotic *B. longum* strains were isolated from both active and non-active IBD patients. This supports the idea that strains with potential probiotic capacities can be isolated even from patients in active disease state. Even if these strains did not manage IBD symptoms when naturally occurring in the IBD intestinal microbiota, they could have benefcial efects on gut health when supplemented in higher amount or in other patients, where other interactions could occur with the host microbiota and immune system. Further pre-clinical and clinical studies could confrm the potential use of IBD isolated *B. longum* strains for applications as probiotics in general gut health and/or in IBD treatment.

Conclusions

We conducted a culturomics approach to characterize the diversity of bifdobacteria and their functionality in the microbiota of active and non-active IBD patients. Relative bifdobacteria abundances were high in *B. dentium* and *B. adolescentis*. *B. dentium* were mainly associated to UC and *B. adolescentis* to CD whereas no specifc bifdobacterial taxonomic profles were highlighted according to active or non-active disease states. Additionally, *B. dentium* relative abundances increased with age, along with a decrease in *B. longum* and *B. pseudocatenulatum*.. Correlations were found between the antipathogenic capacities of the strains and their origin, but overall, we highlighted that bifdobacteria with possible probiotic potential (antipathogenic activity, EPS production and anti-infammatory activity) were isolated from both non-active and active CD and UC patients. We identifed interesting *B. longum* strains for further characterization and potential application as probiotics for human gut health.

Abbreviations

Supplementary Information

The online version contains supplementary material available at [https://doi.](https://doi.org/10.1186/s12866-024-03368-4) [org/10.1186/s12866-024-03368-4](https://doi.org/10.1186/s12866-024-03368-4).

Acknowledgements

We gratefully thank Christel Neut for her contribution to strain isolation in this study. We thank Christel Rousseaux and Intestinal Biotech Development for the collaboration and access to bifdobacteria strains isolated from IBD patients. We thank the Foundation DigestScience for its help in the supervision of the study. We thank the Nutrition, Microbiota and Health Center of Excellence at the Lesafre Institute of Science and Technology for their help setting up the anti-infammatory screening model. We thank Nabil Bosco, Rober Kemperman and Sherrill Cropper for their comments on the draft of this article.

Authors' contributions

BF and CM conceived and designed the project and were major contributors in writing the manuscript. PD performed clinical evaluation of all patients and provided patient data. SB managed the strain collection constituted, performed functional screening, analyzed, and interpreted all data, and was the major contributor in writing the manuscript. LD managed the strain collection. MT identifed all strains by MALDI-TOF and ND, SD, MS and MDSV by WGS. ND set up automated high throughput methods for functional characterization. ED performed immunomodulatory screening. HAA participated in study design and managed collaborations between the diferent parties. FM performed all statistical analysis. All authors read and approved the fnal manuscript.

Funding

This work was funded by Lesafre International.

Availability of data and materials

Data supporting the findings of this study are available within the supplementary fle provided. Associated DNA sequences were deposited in SRA under accession number PRJNA1090283.

Declarations

Ethics approval and consent to participate

Fecal samples were collected during medical care at Lille University Hospital by Pr. Desreumaux during routine gastroenterological consultations and Bacterial strains were further isolated from these samples by IBD Biotech, as secondary use of biological material, as approved by the French law (Article L1211-2). A local ethics committee (Comité de Protection des Personnes Nord Ouest IV, CHRU Lille, France) approved the study (No. DC-2008–642) and all subjects signed clear informed consent.

Consent for publication

Not applicable.

Competing interests

This study was a collaboration between Lesafre International and Infnite (Inserm, CHU Lille, U1286, France). SB, LD, ED, SD, MS, ND, MDSV, FM, HAA and CM are employees of Lesafre International, who funded this study. P.D. reports personal fees from Lesafre and several pharmaceutical companies, Abbott, Amgen, Biocodex, Biofortis, Biogen, Biokuris, Ferring, Fresenius, Janssen Kitozyme, MSD, Norgine, Pfzer, Sandoz, Shire, Takeda, Tillotts, and UCB, outside of the submitted work. Other authors declared no competing interests.

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Received: 14 February 2024 Accepted: 12 June 2024

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