RESEARCH

Genome insights into the pharmaceutical and plant growth promoting features of the novel species Nocardia alni sp. nov

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Abstract

Background: Recent studies highlighted the biosynthetic potential of nocardiae to produce diverse novel natural products comparable to that of Streptomyces, thereby making them an attractive source of new drug leads. Many of the 119 Nocardia validly named species were isolated from natural habitats but little is known about the diversity and the potential of the endophytic nocardiae of root nodule of actinorhizal plants.

Results: The taxonomic status of an actinobacterium strain, designated ncl2T, was established in a genome-based polyphasic study. The strain was Gram-stain-positive, produced substrate and aerial hyphae that fragmented into coccoid and rod-like elements and showed chemotaxonomic properties that were also typical of the genus Nocardia. It formed a distinct branch in the Nocardia 16S rRNA gene tree and was most closely related to the type strains of Nocardia nova (98.6%), Nocardia jiangxiensis (98.4%), Nocardia miyuensis (97.8%) and Nocardia vaccinii (97.7%). A comparison of the draft genome sequence generated for the isolate with the whole genome sequences of its closest phylogenetic neighbours showed that it was most closely related to the N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data. Corresponding taxogenomic data, including those from a pan-genome sequence analysis showed that strain ncl2T was most closely related to N. jiangxiensis, N. miyuensis and N. vaccinii strains, a result underpinned by average nucleotide identity and digital DNA-DNA hybridization data.

Conclusions: Our genome-based taxonomic study showed that isolate ncl2T formed a new centre of evolutionary variation within the genus Nocardia. This novel endophytic strain contained natural product biosynthetic gene clusters predicted to synthesize novel specialised products, notably antibiotics. Genome mining of the N. alni and N. vaccinii strains contained 36 and 29 natural product-biosynthetic gene clusters, respectively, many of which were predicted to encode for a broad range of novel specialised products, notably antibiotics. Genome mining of the N. alni and N. vaccinii strains contained 36 and 29 natural product-biosynthetic gene clusters, respectively, many of which were predicted to encode for a broad range of novel specialised products, notably antibiotics. Genome mining of the N. alni and N. vaccinii strains contained 36 and 29 natural product-biosynthetic gene clusters, respectively, many of which were predicted to encode for a broad range of novel specialised products, notably antibiotics. Genome mining of the N. alni and N. vaccinii strains contained 36 and 29 natural product-biosynthetic gene clusters, respectively, many of which were predicted to encode for a broad range of novel specialised products, notably antibiotics.

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Introduction

The actinobacterial genus *Nocardi*a [1], the type genus of the family *Nocardiaceae* [2] emend. Zhi et al. [3], has a long and convoluted taxonomic history mainly due to an overreliance placed on morphological properties [4, 5]. The application of polyphasic taxonomic procedures led to marked improvements in the classification of nocardiae and related mycolic acid containing actinobacteria [6]. In general, the genus encompasses aerobic, Gram-stain-positive, acid-alcohol-positive, nonmotile, chemoorganotrophic actinobacteria which form rudimentary to extensively branched substrate hyphae that fragment into coccoid to rod-shaped elements, aerial hyphae may only be visible microscopically; the diamino acid of the peptidoglycan is meso-diaminopimelic acid (A2pm), the characteristic whole-organism sugars are arabinose and galactose; diphosphatidylglycerol, phosphatidylethanolamine, phosphatidylglycositol and phosphatidylglycinositol mannosides are the major polar lipids; the fatty acids consist of straight-chain, saturated, unsaturated and 10-methyl (tuberculosteric) components; mycolic acids have 46-64 carbon atoms and up to four double bonds; the predominant respiratory quinone is a hexahydrogenated menaquinone with eight isoprene units where the two end ones are cyclized (MK-8[H6-ω-cyclo]) and the DNA G+C content ranges from 63-72 mol% [5, 7].

Many of the 119 *Nocardi*a species with validly published names (https://www.bacterio.net/) are recognized using combinations of genotypic and phenotypic properties [7–9]. Most of these taxa are composed of strains isolated from natural habitats but the best-known species contain causal agents of serious suppurative and granulomatous diseases in humans and animals, especially mycetoma and nocardiosis [10–12]. In contrast, *Nocardi*a *vaccini* produces galls on blueberry plants [13]. Soil is probably the primary reservoir for *Nocardi*a strains as they are found in diverse soil types, including acidic forest [14, 15], arid [16], Cerrado [17], karst cave [18], rhizosphere [19, 20] and saline soils [21, 22]. However, they have also been isolated from marine habitats, especially from sponges [23, 24], as well as from the gut of fungus-growing termites [25] and are increasingly being isolated from plant tissue [26], notably from nodules of actinorhizal plants suggesting that they may have a role in promoting plant growth and inhibiting phytopathogens [27, 28]. Two *Nocardi*a strains isolated from *Casuarina glauca* nodules induced root nodule-like structures in the original host plant [29].

Nocardiae are an important source of novel antibiotics [30, 31], as exemplified by the production of amicoumacin B from *Nocardi*a *jinanensis* [32], asterobactin from *Nocardi*a *asteroides* [33], brasilcardin A from *Nocardi*a *brasiliensis* [34], nocardicins from *Nocardi*a *uniformis* subsp. *tsuyamanensis* [35] and tubelactomicin A from *Nocardi*a *vinacea* [36]. A comparative survey of nocardial genomes showed that their biosynthetic potential to produce diverse novel natural products is comparable to that of better studied actinobacterial taxa, such as *Amycolatopsis* and *Streptomyces*, thereby making them an attractive source of new drug leads [37]. These researchers showed that *Nocardi*a strains from diverse sources, including clinical material, were equally spread across six phylogenetic clades and found that the genomes of the more pathogenic strains were, on average, slightly smaller than those of most of the other genomes (7.4 Mbp against 7.8 Mbp) and contained fewer BGCs (32.5 against 36.5). Similarly, information from the genome of *Nocardi*a *cyriacigeorgica* shows evidence of adaptation from a saprophytic to a pathogenic lifestyle [38].

The present study was designed to establish the taxonomic status of *Nocardi*a strain ncl2T, isolated from a root nodule of an actinorhizal plant, and to determine its biotechnological and ecological potential. The strain was the subject of a genome-based taxonomic study which showed that it formed a new centre of evolutionary variation within the genus *Nocardi*a, the name proposed for this organism is *Nocardi*a *alni* sp. nov. with isolate ncl2T as the type strain. The genomes of *N. alni* and *N. vaccini* strains contained natural product biosynthetic gene clusters predicted to synthesize novel specialised products, notably antibiotics and genes associated with the expression of plant growth promoting compounds. Statistical comparison between genomic features of the isolate and its taxogenomic neighbours were undertaken to establish any positive correlations between them.

Materials and methods

Isolation, maintenance and cultivation

Strain ncl2T was isolated from a root nodule of an *Alnus glutinosa* plant growing in Leazes Park, Newcastle upon Tyne, UK, as described in Ghodhbane-Gtari et al. [27]. The permission to collect the root nodules was obtained and this study complies with local and national regulations. Sterile lobes of harvested nodules prepared using the procedure described by these workers were placed onto BAP agar plates [39] which were incubated until single actinobacterial colonies were detected. One such
strain was checked for purity and maintained in 35% (w/v) glycerol at -80°C, as was Nocardia vaccinii DSM 43285T obtained from the German Collection of Microorganisms and Cell Cultures (DSMZ). Working cultures of these strains were kept on yeast extract-malt extract agar slopes (International Streptomyces Project [ISP] medium 2) [40]. Biomass for the chemotaxonomic analyses carried out on the strains was harvested from ISP2 broths shaken at 200 rpm in baffled flasks for 7 days at 28°C. The harvested cells were washed in distilled water and freeze dried.

**Chemotaxonomic, cultural and morphological properties**

Smears from ISP2 agar slopes of the isolate were Gram-stained using Hucker's modification [41] and the presence of fragmenting branched hyphae sought by light microscopy. The isolate was examined for its ability to grow over a wide range of temperatures (4°C, 10°C, 15°C, 25°C, 28°C, 37°C, 42°C and 45°C) using ISP2 agar as the basal medium. Growth and cultural properties were recorded on GYM (DSMZ medium 65), nutrient agar (NA), peptone-meat extract-glucose agar (DSMZ medium 250) and tryptic soy agar (TSA) after 7 days at 28°C.

The isolate and N. vaccinii DSM 43285T were examined for chemotaxonomic properties known to be of value in nocardial systematics [5, 25]. Standard chromatographic procedures were used to establish the diamino acid of the wall peptidoglycan [42], whole organism sugars [43] and polar lipid profiles [44]). Cellular fatty acids were extracted and methylated after Miller [45], as modified by Kyukendall et al. [46], and analysed by gas chromatography (Agilent instrument, model 6890N). The resultant peaks were identified using the Standard Micribial Identification (MIDI) system, version 4.5 and the ACTINO6 database [47]. Mycolic acids were extracted using the procedure described by Minnikin and Goodfellow [48], purified and their profiles determined by gas chromatography (Agilent instrument, model 6890N).

**Genome sequencing**

Genomic DNA was extracted from wet biomass of a single colony of isolate ncl2T grown on ISP2 agar for 10 days at 28°C. The extracted DNA was purified and quantified following the protocol of MicrobesNG, Birmingham (UK). Genomic DNA libraries and sequencing were achieved using an Illumina HiSeq instrument and the 250bp paired end protocol, as used in the service provided by MicrobesNG. The draft genome sequence was annotated using the RAST-SEED webserver with default options [49].

**Phylogeny**

An almost full length 16S rRNA gene sequence (1523 bp) extracted directly from the draft genome sequence of isolate ncl2T was deposited in the GenBank databases under accession number MZ014381. The resultant gene sequence was compared with corresponding gene sequences of closely related Nocardia strains retrieved from the EzBioCloud server [50]. Phylogenetic trees based on single 16S rRNA genes and corresponding genome sequences were inferred using the Type Strain Genome Server (TYG), the high throughput Genome to Genome Distance Calculator (GGDC) webserver of Meier-Kolthoff et al. [51]. Average nucleotide identity (ortho ANI, [52]) and digital DNA-DNA hybridization (dDDH) [53] similarities were determined between the isolate and its phylogenomic neighbours using the ANI calculator from the EzBioCloud (http://www.ezbiocloud.nettools/ani) and the GGDC webservers, respectively.

The type strain of Nocardia casuarinae, isolated from root nodules of Casuarina glauca [27] was included for comparative purposes, as was that of Nocardia pseudobrasiliensis which was isolated from a leg abscess of a patient suffering from ulcerative colitis [54].

**Comparative genomic analyses**

The genome sequence of isolate ncl2T was compared with that of N. vaccinii NBRC 15922T (GenBank accession number BDCC00000000.1), its nearest taxogenomic neighbour, and with those of other phylogenomic relatives, namely Nocardia jiangxiensis NBRC 101359T (GenBank accession number BAGB00000000.1), Nocardia miyunensis NBRC 108239T (GenBank accession number BDBQ00000000.1), as well as with N. casuarinae BMG 51109T (GenBank accession number JAFQ00000000) and N. pseudobrasiliensis DSM 44290T (GenBank accession number QQBC00000000.1). These strains were from diverse sources, namely from bud-proliferating galls on blueberry [13], the rhizosphere of goose-grass (*Eleusine indica*), a pine forest soil [14], the root nodule of *Casuarina glauca* and a leg abscess of the patient with ulcerative colitis, respectively. The draft genome assemblies of the strains were annotated using the RAST-SEED webserver [49, 55] with default options. Genome-based species identification was achieved using the TrueBac ID System v1.92, DB:20190603 [https://www.truebacid.com/] [56] and the algorithm proposed by Chun et al. [57]. For the comparative genomic analyses, homologous regions in the target genomes were determined to query ORFs using the USEARCH program version v8.1.1861, and aligned using a pairwise global alignment [58]. The matched region in the subject contig was extracted and saved as a homolog.
A pairwise gene-to-gene comparison of each genome was conducted using USEARCH and the gene contents among the isolates and related strains compared using the reciprocal homology search tool as described in Chun et al. [59] and Ha et al. [60]. Reciprocal relations are determined if two different genes give bidirectionally top hits to one another. A pairwise orthologous group is defined if the pair of genes has reciprocal relations. The term pairwise orthologous groups (POGs) was coined for these collections of reciprocally linked orthologs. After the initial grouping, partial genes that are grouped due to their short sequence length are targeted for clustering analysis against the POGs using UCBLUST (>95 % identity). The coding sequences (CDSs) were classified into groups based on their roles, with reference to orthologous groups (EggNOG 4.5; http://egg nogdb.embl.de) [61].

Statistical comparisons of the genomic properties of the isolate and those of closely related strains (genome size, digital G+C content, number and median length of codon sequences) were carried out to determine possible correlations between them.

Phenotypic properties
Strain ncl2T and N. vaccinii DSM 43285T were screened for a broad range of phenotypic properties, including their ability to metabolize diverse sole carbon and nitrogen sources, to grow in the presence of several concentrations of sodium chloride, at a range of pH values and in the presence of antibiotics using GINII microplates and an OMNILOG device (Biolog Inc., Hayward, CA, USA), as described previously [8]. The resultant data were analysed using version 1.3.36 of the OPM package [62, 63]. They were also tested for their ability to produce niacin, arylsulfatase after 3 days [64], and to reduce tellurite [65]. All of these tests were carried out in duplicate using a standard inoculum. Enzymatic and additional metabolic properties of the strains were determined using API-ZYM kits and the protocol provided by the manufacturer (Biomerieux, France).

Results and discussion
Chemotaxonomic, cultural and micromorphological properties
Isolate ncl2T showed a range of phenotypic properties consistent with its classification in the genus Nocardia [5]. It is aerobic, Gram-strain-positive and forms extensively branched substrate hyphae and aerial hyphae that fragment into coccoid to rod-like elements. Beige-pinkish aerial mycelia were formed on DSMZ 65; ISP2 and TSA agar after 10 days at 28°C and 37°C (Fig. S1). The strain grew well on DSMZ 65 and ISP2 agar at 28°C and 37°C, but not at 4°C, 10°C, 15°C, 25°C, 42°C or 45°C, nor did it grow on nutrient agar. It contained meso-A2pm as the diamino acid of the peptidoglycan, arabinose, galactose and glucose as cellular sugars; mycolic acids with 42-62 carbon atoms; the major fatty acids (>10%) were C16:0 and C18:1 ω9c (Table S1) and the polar lipid pattern consisted of diphosphatidylglycerol (DPG), phosphatidylethanolamine (PE), phosphatidylaminolipid (PI), phosphoglycolipid (PGL), and unidentified phospholipids (PL1-2), aminolipid (AL) and lipids (L1-2) as shown in Fig. S5.

Phylogeny
The phylogenetic tree (Fig. 1) shows that strain ncl2T forms a distinct branch in the Nocardia 16S rRNA gene tree which lies between lineages composed of the type strains of Nocardia nova [66], Nocardia aobensis [67], Nocardia cerradoensis [17], and N. vaccinii, and N. jiangxiensis and N. miyunensis, respectively. The close relationship between the N. jiangxiensis and N. miyunensis strains has been recorded repeatedly [14, 24, 25, 68, 69]. Isolate ncl2T was most closely related to N. nova NBRC 15556T, these strains shared a 16S rRNA gene sequence similarity of 98.6%, which corresponds to 20 nucleotides differences. It also showed relatively low sequence similarities with the N. aobensis (97.2%), N. cerradoensis (97.0%), N. jiangxiensis (98.4%), N. miyunensis (97.8%) and N. vaccinii (97.7%), these relationships were not supported by high bootstrap values (Fig. 1).

Greater confidence can be placed in the topology of phylogenomic tree when compared against corresponding 16S rRNA gene tree as the former is based on millions, not hundreds, of unit characters [70]. The phylogenomic tree (Fig. 2) shows that isolate ncl2T forms a distinct branch that is most closely related to an evolutionary group composed of the type strains of N. jiangxiensis, N. miyunensis and N. vaccinii, a relationship supported by a 100% bootstrap value. The members of this lineage form a subclade next to a well-supported taxon composed of the type strains of N. aobensis, N. cerradoensis and N. nova.

The recommended threshold used to distinguish between closely related strains based on ANI and DDH values are 95% to 96% and 70%, respectively [57, 71]. Table 1 shows that on this basis isolate ncl2T can be distinguished from its closest phylogenomic neighbours and from the type strains of N. casuariinae and N. pseudobrasiliensis. It is also clear that it is most closely related to the N. jiangxiensis, N. miyunensis and N. vaccinii strains although the shared ANI and dDDH similarities are low falling within the range 80.2% to 80.7% and 24.4% to 24.9%, respectively.
Genome features

Key genomic features of the isolate and the associated reference strains are shown in Table 2. The in silico G+C content of the strains fall within a narrow range, namely 66.7 to 68.9%. In contrast, the corresponding genomes show more variation ranging from 8.4 to 10.5 Mbp in size. The isolate is most closely related to N. vaccinii DSM 43285\textsuperscript{T}, as these strains have genomes of a similar size (9.9 Mbp and 9.2 Mbp, respectively), very similar dG+C values (67.0% and 66.7%, respectively), and a similar number of CDCs (8969 and 8502) and tRNA genes (49). In contrast, the N. jiangxiensis and N. miyunensis strains have larger genomes, namely 10.45 and 10.52 Mbp, respectively, and a similar, albeit larger, number of CDS;
these results underpin the close relationship between these strains as found in 16S rRNA gene sequence analyses. The *N. pseudobrasiliensis* strain, a representative of a species associated with invasive human diseases [54], has the smallest genome, thereby providing further evidence that genome sizes of clinically significant nocardiae are lower than corresponding results from non-pathogenic strains [37]. It is also interesting that the type strain of *N. casuarinae*, which induces root nodule formation in *C. glauca*, has a genome size similar to that of *N. pseudobrasiliensis* DSM 44290T.

A comparison of the taxogenomic features (genome size, digital GC content, number of CDSs, median length of CDS) of strain ncl2T and the five associated reference strains shows that the number of CDSs is positively correlated with genome size and with the coefficient of determination (R2=0.94), this means that 94% of the data points support the predicted regression lines (y=c[176]+c[896]x), shown in Fig. S2. The frequency plot of the pan genome orthologous groups (POGs) of the strains highlight POGs involved in amino acid and carbohydrate metabolism, information storage and processing (eg. recombination and replication) and cellular processes and signaling (Fig. S3). However, no clear correlation was found between genome size and the number of orthologous gene groups though the frequency plots of the POGs for the genomes of the *N. casuarinae*, *N. jiangxiensis*, *N. miyunensis* strains were similar with little evidence of quantitative variation.

Comparison of the functional categories between genes in the core and pan genomes using COG/EggNog software gives results in good agreement with those found in the SEED analysis. Apart from genes with unknown function and unassigned categories, the core genomes are composed mainly of genes involved in amino acid transport and metabolism, energy production, and conversion and transcription, as shown in Fig. S4a. However; the pan genome of the strains also contain genes associated with carbohydrate metabolism, clustering based systems and the metabolism of amino acids and derivatives, as presented in Fig. S4b. Around 75% of the core genomes are composed of genes assigned to defined categories whereas less than 40% of those in the pan genomes are associated with functional categories based on COG and SEED pathways. When the strains were examined for strain specific CDSs the highest number were for strain ncl2T with 2162 and the *N. pseudobrasiliensis* DSM 44290T with 2187. The highest similarity was between the pan genome of strain ncl2T and its closest taxogenomic neighbour, *N. vaccinii*, then by the *N. jiangxiensis*, *N. miyunensis* strains which were isolated from acidic soil; no correlation was found between the genome gene contents of the strains and the habitats from which they were isolated.

**Phenotypic properties**

Identical results were obtained for nearly all of the duplicated cultures included in the phenotypic tests. Table 3 shows that the isolate can be distinguished from *N. vaccinii* DSM 43285T as it produced niacin, utilized L-alanine and L-serine and an extensive range of organic compounds, including β-gentiobiose, dextrin, D-maltose, D-mannose, pectin, D-raffinose, L-rhamnose, sucrose, D-trehalose and D-turanose, was resistant to fusidic acid, lincomycin, minocycline, troleandomycin and

### Table 1: Average nucleotide identity (ANI) and digital DNA-DNA hybridization (dDDH) values between strain ncl2T and its closest phylogenomic neighbours and with the type strains of *N. casuarinae* and *Nocardia pseudobrasiliensis*.

| Species            | Strain designation | dDDH values | ANI values |
|--------------------|--------------------|-------------|------------|
| *N. aobensis*      | NBRC 100429T       | 22.4        | 77.9       |
| *N. casuarinae*    | BMG 51109T         | 23.0        | 78.9       |
| *N. cerradoensis*  | NBRC 101014T       | 22.4        | 77.8       |
| *N. jiangxiensis*  | NBRC 101359T       | 24.6        | 80.6       |
| *N. miyunensis*    | NBRC 108239T       | 24.9        | 80.7       |
| *N. nova*          | NBRC 15556T        | 22.4        | 77.9       |
| *N. pseudobrasiliensis* | DSM 44290T    | 22.4        | 78.7       |
| *N. vaccinii*      | NBRC 15922T        | 24.4        | 80.2       |

### Table 2: Genomic features of isolate ncl2T (1) and the type strains of *N. vaccinii* (2), *N. miyunensis* (3), *N. jiangxiensis* (4), its closest phylogenetic relatives, and with those of *Nocardia casuarinae* (5) and *N. pseudobrasiliensis* (6).

| Genomic features | 1     | 2     | 3     | 4     | 5     | 6     |
|------------------|-------|-------|-------|-------|-------|-------|
| Genome size (Mbp) | 9.9   | 9.2   | 10.5  | 10.4  | 8.8   | 8.4   |
| Digital G+C content | 67.0  | 66.7  | 66.9  | 66.7  | 68.9  | 67.3  |
| No. of rRNA genes | 12    | 4     | 4     | 3     | 9     | 8     |
| No. of tRNA genes | 49    | 49    | 49    | 49    | 56    | 51    |
| No. of CDSs      | 8969  | 8502  | 9567  | 9695  | 7746  | 7949  |
| Median length of CDS (bp) | 831   | 816   | 831   | 822   | 819   | 822   |
vancomycin, and grew in the presence of lithium chloride, guanidine hydrochloride, sodium bromate, sodium chloride (4% and 8%), tetrazolium blue, and tetrazolium violet. In contrast, only the N. vaccinii strain utilized α-keto-glutaric acid, L-malic acid, mucic acid, propionic acid and bromo-succinic acid, it also showed greater activity in the API-ZYM tests.

Strains ncl2T and N. vaccinii NBRC 15922T can also be distinguished using chemotaxonomic and genomic traits. They showed qualitative and quantitative differences in fatty acid profiles, as exemplified by the presence of a trace of C17:1ω9c and a high proportion of C18:010-methyl in the N. vaccinii strain and a higher proportion of C18:1ω9c in the isolate (24.4% against 9.6%) (Table S1). These properties distinguish them from the type strains of N. jiangxiensis and N. miyunensis as shown in Table S1. Further, only the isolate contains glucose in whole cell hydrolysates and unidentified aminolipid and glycolipid in its polar lipid profile, unlike N. vaccinii DSM 43285T (Fig. S5). The isolate contains mycolic acids ranging from 42 to 62 carbons with predominant amounts of C42 and C44 whereas the N. vaccinii strain has mycolic acids ranging from 52 to 62 carbons with C54, C56, and C58 as major components. Both strains have meso-A2pm

### Table 3 Phenotypic, enzymatic and biochemical properties that distinguish strain ncl2T from Nocardia vaccinii DSM 43285T

| Characteristics | Isolate ncl2T | N. vaccinii DSM 43285T |
|-----------------|---------------|------------------------|
| **GEN III Biolog microplate tests** |               |                        |
| **Sugars:**     |               |                        |
| D-Cellobiose, D-galactose, β-gentiobiose, D-maltose, D-mannose, D-raffinose, L-rhamnose, sucrose, D-trehalose, and D-turanose | + | - |
| **Polymers:**   |               |                        |
| Dextrin, pectin |               |                        |
| **Tolerance to:** |               |                        |
| Lithium chloride, guanidine hydrochloride, sodium bromate, sodium chloride (4% and 8%), tetrazolium blue, and tetrazolium violet | + | - |
| **Sugar alcohols:** |               |                        |
| D-Arabitol      |               |                        |
| D-Mannitol      |               | +                      |
| **Oxidation of amino acids:** |               |                        |
| L-Alanine, L-serine |               | +                      |
| L-Aspartic acid |               | +                      |
| **Resistant to:** |               |                        |
| Fusidic acid, lincomycin, minocycline, troleandomycin, and vancomycin | + | - |
| **Sugar acids:** |               |                        |
| L-Galactonic acid-γ-lactone, andD-galacturonic acid | + | - |
| **Organic acids** |               |                        |
| γ-amino-γ-Butyric acid, α-hydroxy-γ-butyric acid, L-lactic acid, D-lactic acid methyl ester, D-p-hydroxy-phenylacetic acid, methyl pyruvate, and quinic acid, α-keto-Glutamic acid, L-malic acid, mucic acid, propionic acid, and bromo-succinic acid, | + | - |
| **API ZYM tests:** |               |                        |
| Alkaline phosphatase, α-chymotrypsin, cystine aminopeptidase, esterase (C4), esterase lipase (C8), lipase (C14), trypsin, and valine aminopeptidase | - | + |
| **Other tests:** |               |                        |
| Niacin |               | +                      |

+ , positive result; - , negative result

Both strains reduced potassium tellurite and gave positive results for acid phosphatase, β-galactosidase α-glucosidase; oxidized, D-glucose, D-glucic acid, glycerol, and myo-inositol (sugars); acetic acid, acetacetic acid, butyric acid, β-hydroxy-butyric acid, α-keto-butyric acid, citric acid, and D-saccharic acid, (organic acids); L-arginine, L-glutamic acid and D-serine #2 (amino acids); were resistant to nalidixic acid, and rifamycin sv (antibiotics); and grew in presence of potassium tellurite, and Tween 40 (inhibitory compounds); and in the presence 1% sodium lactate, sodium chloride (1%) (salts) and at pH 5–6. The strains gave negative results for arylsulfatase after 3 days, D-fucose, L-fucose, D-fructose, D-fructose-6-phosphate, N-acetyl-D-galactosamine, N-acety-D-glucosamine, D-glucose-6-phosphate, glucuronamide, 3-O-methyl-D-glucose, β-methyl-D-glucoside, α-D-lactose, D-melibiose, D-salicin, D-sorbitol, and D-stachyose (sugars); D-glucic acid, D-malic acid, N-acetyl-neuraminic acid, and L-γ-pyrogulutaric acid (organic acids), D-aspartic acid, glycine-proline, L-histidine, and D-serine #1 (amino acids); inosine (nucleoside); aztreonam (antibiotic); gelatin (polymer), niaproof, and sodium formate (inhibitory compounds); α-fucosidase; α-galactosidase, N-acetyl-β-glucosaminidase; β-glucuronidase, and α-mannosidase (API-ZYM tests)
as the diamino acid of the wall peptidoglycan, arabinose and galactose as diagnostic whole cell sugars and C\textsubscript{16:0} as the major fatty acids. Furthermore, the genome size of the isolate is larger than that of the \textit{N. vaccinii} strain (9.9 against 9.2 Mbp).

**Specialized metabolite biosynthetic gene clusters**

Antismash 5.0 [72] predicts NP-BGCs based on the percentage of genes from the closest known bioclusters which show BLAST hits to the genomes of the strains under consideration. The genomes of strain ncl2\textsuperscript{T} and \textit{N. vaccinii} NBRC 15922\textsuperscript{T} contained 36 and 29 well-defined bioclusters that are predicted to encode for a broad range of specialized metabolites albeit with low levels of gene identity, as shown in Table S2. The genomes of the strains are well equipped to synthesize non-ribosomal peptide synthetases, type I polyketides, ribosomally synthesized and post-translationally modified peptides, as well as betalactone (3% gene similarity) and carotenoid-like terpene (18% gene similarity) compounds. They have the genetic capacity to produce products most closely related to himastatin (3% gene similarity), an antitumor antibiotic produced by \textit{Streptomyces hygroscopicus} [73], steffimycin D (8% gene similarity), which was initially produced by a \textit{Streptomyces} strain and inhibits ras-oncogen expressed cells [74], and teicoplanin, a product of a \textit{Streptomyces} strain that inhibits growth of Gram-positive bacteria, including \textit{Enterococcus faecalis} and methicillin-resistant \textit{Staphylococcus aureus} (MRSA) strains [75]. The strains also contain bioclusters predicted to synthesize arylpene-like compounds that are structurally and functionally similar to carotenoids [76] and which show antimicrobial and antioxidant activity [77]. They also have bioclusters predicted to encode for ectoine (100% gene similarity), a protective molecule which enables bacteria to survive extreme conditions [78].

The genome of strain ncl2\textsuperscript{T}, unlike that of \textit{N. vaccinii} NBRC 15922\textsuperscript{T}, contains presumptive NP-BGCs associated with the synthesis of amycalamycin A and B (2% gene similarity), type I polyketides that are cytotoxic for breast cancer cell lines [79], pepticinnamin E (6% gene similarity) which has the potential to treat cancer and malaria [80] and echosides A, B, C, D and E (11% gene similarity) and stambomycin A to D (36% gene similarity), which are antitumor antibiotics [81]. Strain ncl2\textsuperscript{T} also contains bioclusters predicted to encode for the antitubercular polyketides atratumycin (7% gene similarity) and capreomycin A and B (6% gene similarity), which are produced by \textit{Streptomyces aratus} and \textit{Streptomyces capreoli}, respectively [32, 82]. Additional presumptive bioclusters are linked to the synthesis of lasolocid (3% gene similarity), a betalactone produced by \textit{Streptomyces lasalliensis}, which has antibacterial and coccidiostatic properties [83] and tiacumicin B (3% gene similarity), a type I polyketide active against nosocomial diarrhea caused by \textit{Clostridium difficile} [84]. Other bioclusters are predicted to encode for the antifungal agents fengycin (3% gene similarity) and nystatin (31% gene similarity) that are produced by \textit{Bacillus subtilis} and \textit{Streptomyces noursei} strains, respectively [85–87].

The genome of the \textit{N. vaccinii} strain harbours several bioclusters absent from that of strain ncl2\textsuperscript{T}, including ones predicted to encode for polyketides (Table S2). These bioclusters include cyphomycin (2% gene similarity), which is produced by a \textit{Streptomyces} strain and is used to control multidrug resistant fungal pathogens [88] and caniferolides A-D, that are synthesized by \textit{Streptomyces caniferus} and inhibit the growth of \textit{Candida albicans} and \textit{Aspergillus} spp. [89]; caniferolide A has been used to treat Alzheimer’s disease [90]. Further, the \textit{N. vaccinii} strain has the genetic capacity to synthesise cremimycin MJ635-865F and tetranyrimycins A and B, these antibiotics are produced by \textit{Streptomyces} strains and show activity against MRSA strains [91, 92]. In addition, the strain contains a presumptive biocluster associated with the expression of leinamycin (2% gene similarity), a betalactone terpene produced by \textit{Streptomyces atrolo-vaceus} which shows antibacterial and antitumor activity [93].

It can be concluded that strain ncl2\textsuperscript{T} and \textit{N. vaccinii} NBRC 15928\textsuperscript{T} have genomes rich in NP-BGCs, notably ones predicted to express for putatively novel polyketide and non-ribosomal peptide compounds thereby providing further evidence that nocardiae are a potentially prolific source of new bioactive compounds [37]. It is particularly interesting that these strains have the capacity to synthesise antifungal and antibiotics given their association with plant tissues. Clearly, nocardiae should feature more prominently in natural product discovery campaigns.

**Plant growth promoting properties**

Comparative genome mining of strain ncl2\textsuperscript{T} and the type strains of \textit{N. jiangxiensis}, \textit{N. miyunensis} and \textit{N. vaccinii}, its closest phylogenomic neighbours, revealed the presence of genes associated with direct (eg. phosphate solubilization, phytohormone production) and indirect (eg. lytic enzyme and siderophore production) mechanisms that promote plant growth. \textit{Nocardia casuarinae} BMG51109\textsuperscript{T} and \textit{N. pseudobrasiliensis} DSM 44290\textsuperscript{T} were included in these analyses to represent taxa isolated from plant and clinical sources, respectively [27, 54].

Microbes have a pivotal role in making phosphorus available to plants [94] either enzymatically [95] or by producing organic acids and siderophores and other molecules that solubilize inorganic phosphate [96, 97].
The genome of all of the strains, apart from that of the \( N. \) \textit{pseudobrasiliensis} DSM 44290\(^T \), contained genes associated with phosphate regulation and metabolism (Table S3). These included gene \( ppx-gppA \), which is responsible for the solubilization of inorganic polyphosphate [98] and gene \( \text{pstS} \) which encodes for phosphate binding protein \( \text{PstS} \) that is involved in the production of the phosphate ABC transporter [99]. The \( \text{pstS} \) gene was not detected in the genome of the clinical isolate thereby suggesting a possible correlation between the environmental origin of the other strains, namely soil and plant tissues, and phosphate metabolism. The genome of all of the strains contained gene \( \text{senX3} \) which is associated with the production of histidine kinase, a high affinity phosphate transporter which has a role in controlling the phosphate regulon [100].

Phytohormones have a central role in plant growth [101], notably indole-3-acetic acid (IAA) and ethylene; the levels of these and other auxins in plants can be regulated by soil microorganisms able to synthesize them [102]. The genome of all of the strains contained genes encoding for indole-3-glycerol phosphate synthase, the precursor of IAA in the tryptophan biosynthetic pathway in plants [103]. They also contained genes encoding for other components of this pathway, including anthranilate phosphoribosyl transferase (\( \text{trpD} \)), anthranilate synthase (\( \text{trpE} \)), and aminase (\( \text{trp A and B} \) ) [104]. Similarly, gene \( \text{trpF} \), which is associated with the synthesis of anthranilate phosphoribosyl transferase, was present in the genomes of all of the strains, apart from \( N. \) \textit{pseudobrasiliensis} DSM 44290\(^T \). Genes \( \text{pxd1} \) and \( \text{aad} \), which encode for pyridoxine 4-dehydrogenase and aryl-alcohol dehydrogenase (NADP (+)) and are involved in auxin signaling pathways, were found in the genome of strain \( \text{ncl2} \)^\(^T \), \( N. \) \textit{jiangxiensis} NBRC 101359\(^T \), \( N. \) \textit{miyuensis} NBRC 108239\(^T \) and \( N. \) \textit{casurinae} BMG51109\(^T \) (Table S3).

In contrast, the genomes of all of the strains contained genes associated with tricarboxylic acid biosynthesis, as shown in Table S3. However, only the genome of strain \( \text{ncl2} \)^\(^T \) contains gene \( \text{acc} \) that encodes for 1-aminocyclopropane-1-carboxylatedeaminase, an ACC deaminase which reduces toxicity due to high levels of ethylene in plants caused by plant growth promoting rhizobacteria. This enzyme also regulates ethylene levels produced by the plant by converting ACC to ammonia and \( \alpha \)-ketobutyrate [105, 106].

Plant growth promoting microorganisms can also enhance plant growth by modulating biotic stress as they can decrease, neutralize or prevent infections caused by phytopathogens by synthesizing antibiotics and lytic enzymes [107]. The genomes of all of the strains were equipped with genes associated with the production of chitinases and glucoamylases, as shown in Table S3. They also contained genes involved in the biosynthesis of antibiotics, as exemplified by \( \text{fabG} \), \( \text{bacC2} \) and \( \text{hdhA} \) which express for 3-oxoacyl-\( \text{acyl-carrier-protein} \) reductase, bacitracin synthase and 7-alpha-hydroxysteroid dehydrogenase which play a role in the biosynthesis of pentenolactone, bacitracin and clavulanic acid, respectively [108–110]. Further, the genomes of all of the strains, apart from \( N. \) \textit{pseudobrasiliensis} DSM 44290\(^T \), contained gene \( \text{auaJ} \) which encodes for the epoxidase \( \text{LasC} \) that is involved in the synthesis of lasalocid, a polyether antibiotic [111]. In contrast, only strain \( \text{ncl2} \)^\(^T \) contained gene \( \text{tcnO} \) which expresses tetracyclomycin polyketide synthesis 8-O-methyl transferase, a gene is associated with tetracyclomycin biosynthesis [112].

It can be concluded that while strain \( \text{ncl2} \)^\(^T \) is most closely related to the type strains of \( N. \) \textit{jiangxiensis}, \( N. \) \textit{miyuensis} and \( N. \) \textit{vaccinii}, it can be distinguished from them as it forms a distinct branch in the phylogenic tree, has a distinct fatty acid profile and shares low ANI and dDDH values with them. Genomic features, notably genome size and CDS numbers, show that the strain is most closely related to \( N. \) \textit{vaccinii} NBRC15992\(^T \), but can be distinguished from the latter by a wealth of chemotaxonomic, genomic and phenotypic data. It is, therefore, proposed that strain \( \text{ncl2} \)^\(^T \) should be recognized as a new species within the genus \( \text{Nocardia} \) for which the name \( \text{Nocardia alni} \) sp. nov. is proposed.

**Description of \( \text{Nocardia alni} \) sp. nov.**

\( \text{Nocardia alni} \) (a'ni L. gen. fem. n. \( \text{alni} \), of \( \text{Alnus} \) (a genus name), referring to the source of the strain, a root nodule of \( \text{Alnus glutinosa} \))

Aerobic, Gram-stain-positive, nonmotile actinobacterium that forms an extensively branched substrate mycelium and aerial hyphae which fragment into coccosid to rod-like elements. Beige pink aerial hyphae are formed on DSMZ 65, yeast extract-malt extract and tryptic soy agar. Grows from pH 5-7.5 (optimally at pH 7), from 28 to 37°C and in the presence of up to 8% w/v sodium chloride. Produces niacin, reduces potassium tellurite, but is arylsulfatase negative after 3 days. Additional phenotypic properties are shown in Table 3. The diaminoc acid of the peptidoglycan is \( \text{meso-A} \) pm, the whole cell sugars are arabinose, galactose and glucose and the predominant fatty acids are \( \text{C}_{16:0} \) and \( \text{C}_{18:1} \omega9c \). Mycolic acids have 42 to 62 carbon atoms and the polar lipids are diphosphatidylglycerol, phosphatidylethanolamine and phosphatidylinositol, unidentified phosphoglycolipid, phospholipids, lipids, an aminolipid and a glycolipid. The genome size is 9.93 Mbp and the \( \text{in-silico} \) G+C content 67.0%. The genome is rich in biosynthetic gene clusters predicted to encode for new specialised metabolites, notably antibiotics, and with...
genes with the capacity to produce products that promote plant growth.

The type strain ncl2T (DSM 110931T = CECT 30122T) was isolated from a root nodule of *Alnus glutinosa* growing in Leazes Park, Newcastle upon Tyne, UK. The GenBank accession number for the 16S rRNA gene and whole genome sequence of the strain were MZO14381 and JAGPOX000000000, respectively.

**Conclusions**

Novel endophytic nocardiae are being isolated from rhizospheric soil [19, 113], plant roots and stems [26] and from nodules of actinorhizal plants [28–30] as in the case of *N. alni*. Nodular tissues are rich in carbohydrates hence they are excellent habitats for bacteria, including actinobacteria [114, 115]. Filamentous actinobacteria are associated with actinorhizal and legume root nodules, notably novel *Micromonospora* species [116]. The present study suggests that nocardiae, like micromonosporae, have the potential to promote plant growth though eco-physiological studies are needed to establish their interactions with plants, notably their role in root nodules of actinorhizal plants.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12864-021-08257-y.

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**Authors’ contributions**

IN conceived and designed the study. IN, SMH, IB, JC, MG were responsible for analysing the data and preparing the draft paper. All authors read and approved the manuscript.

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**Availability of data and materials**

The 16S rRNA gene and whole genome sequence of strain ncl2T are available in the GenBank under accession numbers MZO14381 and JAGPOX000000000, respectively.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no conflict of interest.

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