



Article The Proteogenome of Symbiotic Frankia alni in Alnus glutinosa Nodules

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Abstract: Omics are the most promising approaches to investigate microbes for which no genetic tools exist such as the nitrogen-fixing symbiotic *Frankia*. A proteogenomic analysis of symbiotic *Frankia alni* was done by comparing those proteins more and less abundant in *Alnus glutinosa* nodules relative to N-replete pure cultures with propionate as the carbon source and ammonium as the nitrogen-source. There were 250 proteins that were significantly overabundant in nodules at a fold change (FC) \geq 2 threshold, and 1429 with the same characteristics in in vitro nitrogen-replete pure culture. Nitrogenase, SuF (Fe–Su biogenesis) and hopanoid lipids synthesis determinants were the most overabundant proteins in symbiosis. Nitrogenase was found to constitute 3% of all *Frankia* proteins in nodules. Sod (superoxide dismutase) was overabundant, indicating a continued oxidative stress, while Kats (catalase) were not. Several transporters were overabundant including one for dicarboxylates and one for branched amino acids. The present results confirm the centrality of nitrogenase in the actinorhizal symbiosis.

Keywords: omics; nitrogenase; hopanoids; actinorhizae

1. Introduction

Frankia is a genus containing soil actinobacteria that establish a nitrogen-fixing root nodule symbiosis with 24 genera and hundreds of species of dicotyledons [1]. The best known of these is *Alnus glutinosa* L., the designated type, which is representative of the genus both for morphological characters and evolutionary rate [2].

Hundreds of *Frankia* strains have been isolated from nodules of the different species and classified according to their physiology [3], host specificity [4] or phylogeny [5]. More recently, many species have been described—the first of which was *Frankia alni* [6]—that can nodulate *Alnus* spp. and *Myrica* spp. Its genome has been deciphered and compared to other species [5] and found to contain no symbiosis island, with all genes related to symbiosis present in several unlinked clusters [7].

This symbiosis between *F. alni* and *Alnus* spp. is initiated by Ca-spiking [8], followed by the synthesis of a root-hair-deforming factor [9] that allows the deformation, entrapment of hyphae and penetration into the cortical cells [10]; the formation of a pre-nodule and the emergence of a mature nitrogen-fixing nodule. On the plant side, the mechanisms involved appear similar to those present in legumes, with several elements of the common symbiotic cascade involved, such as the nuclear transcription factor NIN (nodule inception) [11] and the SYMR (symbiosis) kinase [12]. The genomics of symbiotic and non-symbiotic plant phylogenetic neighbors has shown that the common symbiotic determinants were present in all leguminous and actinorhizal lineages and that loss of symbiotic capacity was accompanied by the loss of *NIN* and *RPG* (rhizobium polar growth) [13].



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On the microbial side, less is known since no genetic tools could be developed despite several attempts, hampering mutagenesis and complementation studies. The genomes revealed no canonical nod genes [5] except in two cl2 strains [14,15] and one cl3 strain [16] where two gene clusters are found, one with $nodAB_1$ and the other with $nodB_2CIJ$; however, these were not detected by proteomics of F. soli exposed to root exudates from compatible *Elaeagnus angustifolia* and incompatible hosts [17] even though the mRNAs had been detected in Datisca nodules [14]. Transcriptomics of Frankia in Alnus nodules [7] has identified nif (nitrogenase), hop (hopanoid synthesis), suf (FeS cluster synthesis) and hup (hydrogenase uptake) as upregulated but has shed no light on the molecular dialogue between the two partners. The determinants and structure of the root deforming factor [10], thus, remain elusive. We also know through omics [18] of early rhizospheric interactions that a cellulase/cellulose synthase cluster is upregulated even though *F. alni* cannot grow using glucose as a carbon source, which is evocative of a local weakening of the hair cell wall and a hardening of the hyphal tip to facilitate entry into root tissues. We also know that the auxin PAA is synthesized in nodules and in pure culture—an auxin that when applied onto roots at 10⁻⁵ M causes emergence of stunted swollen secondary roots that are similar to nodules [19]. It is known that the plant synthesizes peptides that bind to and modify the porosity of vesicles [20] and a non-comparative survey of field nodules has shown nitrogenase, the protein that reduces dinitrogen into ammonium and tricarboxylate cycle (TCA) proteins to be abundant [21].

Other in vitro proteomic studies have been done of *Frankia* with/without ammonium [22] or under osmotic stress [23] and have shown TCA involvement, stress determinants and various regulators.

We undertook the present study to characterize the proteomic response of *Frankia alni* as it forms mature 21 dpi nodules on *A. glutinosa* roots.

2. Materials and Methods

2.1. Plant and Bacterial Material

Frankia alni strain ACN14a [24] cells were maintained in BAP medium [25] with 5 mM propionate as the carbon source and 5 mM NH_4^+ as the nitrogen source buffered to pH 6.5.

Alnus glutinosa seeds were harvested from a tree growing in Lyon, France, used previously [7]. They were grown as before with some modifications: seedlings were transferred to Fahraeus's solution [26] in opaque plastic pots (eight seedlings/pot) and grown for four weeks with $0.5 \text{ g} \cdot \text{L}^{-1}$ KNO₃, followed by one week without KNO₃ before inoculation with *F. alni* [24]. To inoculate seedlings, *Frankia* cells were grown in BAP-PCM medium (4 × 250 mL) until log-phase [27]. Cells were collected by centrifugation, washed twice with sterile ultra-pure water and resuspended in 500 mL of Fahraeus's solution without KNO₃. Cell cultures were homogenized by syringing though a 21G needle. The *Frankia* cell suspensions were applied onto plant roots (symbiotic condition). After 21 days, mature nodules were harvested and ground in liquid nitrogen.

As a reference, *F. alni* cells were inoculated after syringing with a series of needles (21G, 23G, 25G, 27G) and grown for 10 days (corresponding to the end of the exponential phase) in 250 mL of BAP medium with ammonium (5 mM) in agitated 500 mL Erlenmeyer flasks [25] buffered to pH 6.5. No vesicles could be found.

2.2. Proteome Characterization

Each sample was dissolved in LDS 1X buffer (Invitrogen, Carlsbad, CA, USA) with 100 μ L of LDS 1X per 30 mg of pellet. The solutions were warmed at 99 °C for 5 min and subjected to sonication in an ultrasonic bath for 5 min. Each sample was transferred into a 2 mL Precellys (Bertin Technologies, Montigny-le-Bretonneux, France) tube containing 200 mg of glass beads and subjected to three cycles of grinding for 20 s, followed by 30 s pauses. Samples were centrifuged for 40 s at 16,000 × g. The resulting supernatants were transferred into Eppendorf tubes and heated for 10 min at 99 °C. Samples were subjected to a short SDS-PAGE migration and processed as previously described [28]. Briefly, the

whole proteome was extracted as a single polyacrylamide band, reduced with dithiothreitol, treated with iodoacetamide and proteolyzed with sequencing-grade trypsin (Roche, Basel, Switzerland) in the presence of 0.01% of proteaseMAX detergent (Promega, Madison, WI, USA). The resulting peptides were analyzed with an ESI-Q Exactive HF mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) coupled to an Ultimate 3000 176 RSL Nano LC System (Thermo). A volume of 10 μ L of peptides was injected onto a reverse-phase Acclaim PepMap 100 C18 column (3 μ m, 100 Å, 75 μ m id \times 500 mm) and resolved at a flow rate of 0.2 μ L/min with a 60 min gradient of CH₃CN (2.5% to 40%) in the presence of 0.1% HCOOH. The tandem mass spectrometer was operated with a Top20 strategy in data-dependent mode. Only peptide molecular ions with double or triple positive charges were selected for fragmentation with a dynamic exclusion of 10 s as previously described [29]. Tandem mass spectrometry spectra were interpreted using the MASCOT 2.2.04 software (Matrix Science, London, UK) with standard parameters. Proteins were quantified based on their spectral counts, and their abundances were normalized with total spectral counts of all proteins identified as belonging to Frankia for comparing distinct conditions. Proteome comparison between conditions was done according to the TFold module from the PatternLab software (www.patternlabforproteomics.org/; last access 7 January 2022).

2.3. Proteome Data

The mass spectrometry proteomics data were deposited in the ProteomeXchange Consortium via the PRIDE partner repository (www.ebi.ac.uk/pride/archive/; last access 16 December 2021) with the dataset identifier PXD030468 and Project DOI 10.6019/PXD030468.

3. Results

The three biological replicates of symbiotic *Frankia alni* overproduced at a fold change of \geq 2250 proteins (Supplementary Table S1) using a nitrogen-replete propionate-fed pure culture as reference, of which 100 had an FC \geq 4.38 (Table 1). These 100 proteins signing for the specificity of symbiotic state account for 17.9% of the total *Frankia* proteins in nodules, based on their cumulated normalized spectral abundance factors. Conversely, there were 1489 under-produced proteins at an FC \leq 0.5. NifH (FRAAL6813) was the most overabundant protein in this condition with an FC of 291.3.

Table 1. List of the 100 most over-abundant *Frankia* proteins in nodules with the NCBI accession, the FRAAL Id, gene names, the protein name, the fold change, the COG (cluster of orthologous genes), the phylogenetic clusters distribution and the NSAF (normalized spectral abundance factors). The COGs are according to [30]. The distribution code A is present in all *Frankia* and in other actinomycetes, F is present in all *Frankia* strains, f is present in some *Frankia* strains and S is present in symbiotic *Frankia* strains (C1, C1c, C2 and C3 but not in C4). C1, C1c, C2, C3 and C4 indicate it is present in this cluster based on a threshold of 35% Id as seen on Mage [31]. Hypothetical genes have been removed from the list; they are listed in Table S1.

NCBI Accession	FRAAL #	Gene	Functional Description	Fold Change	COG	Distribution	% NSAF Nodule
WP_009741523.1	FRAAL6813	nifH	Nitrogenase iron protein NifH	291.33	Р	S	0.009
WP_011607845.1	FRAAL6811	nifK	Nitrogenase molybdenum–iron protein subunit beta NifK	279.00	С	S	0.005
WP_011607832.1	FRAAL6798	korB	2-Oxoglutarate ferredoxin oxidoreductase subunit beta KorB	91.50	С	S	0.002
WP_011606600.1	FRAAL5516		ATPase	89.00	0	C1	0.003
WP_011607846.1	FRAAL6812	nifD	Nitrogenase molybdenum–iron protein subunit alpha NifD	86.17	С	S	0.002
WP_041939983.1	FRAAL6802	nifU	Iron-sulfur cluster assembly accessory protein NifU	84.17	0	S	0.004

NCBI Accession	FRAAL #	Gene	Functional Description	Fold Change	COG	Distribution	% NSA Nodule
WP_041939982.1	FRAAL6799	korA	2-Oxoglutarate ferredoxin oxidoreductase subunit alpha KorA	82.00	С	S	0.001
WP_011602860.1	FRAAL1673	mopB	Molybdenum-binding protein MopB	64.17	R	S	0.004
WP_011605156.1	FRAAL4025	, ddnB	F420H(2)-dependent quinone nitroreductase, DdnB	44.33	J	S	0.003
WP_011607839.1	FRAAL6805	nifW	Nitrogenase-stabilizing/protective protein NifW	40.33	Р	S	0.003
WP_041939985.1	FRAAL6808	nifX	Nitrogen fixation protein NifX	33.83	-	S	0.002
WP_011602101.1	FRAAL0868	tetR	TetR family transcriptional regulator	33.09	Κ	C1	0.002
WP_011607841.1	FRAAL6807	nif	Associated nitrogen fixation protein NifX	23.83	-	S	0.003
WP_011604521.1	FRAAL3374	mer	Coenzyme F420-dependent N(5),N(10)- methylenetetrahydromethanopterin reductase	22.83	С	f	0.001
WP_011604385.1	FRAAL3236	ddn	Deazaflavin-dependent nitroreductase	18.83	-	C1	0.001
VP_011601860.1	FRAAL0611	ybaB	Nucleoid-associated protein	17.67	0	А	0.001
VP_011603220.1	FRAAL2049		Glycosyl transferase	15.50	М	C1c, C2	0.000
VP_011601350.1	FRAAL0082	arfA	Ammonia release factor ArfA	13.33	М	f	0.001
VP_011606048.1		fadA	3-Ketoacyl-CoA thiolase	12.91	Ι	А	0.003
VP_011602036.1		thiC	Phosphomethylpyrimidine synthase	12.80	Н	А	0.000
VP_041939400.1	FRAAL3855	fdxI	4Fe-4S iron-sulfur binding ferredoxin	12.67	С	А	0.001
VP_011601559.1	FRAAL0301		Transcriptional regulator	12.38	Т	c1	0.000
<pre>WP_011603772.1</pre> WP_011607074.1		gad	Acetyl-CoA acetyltransferase Gamma-aminobutyraldehyde	11.83 11.56	I C	A A	0.000
VP_011604927.1		zuu	dehydrogenase Thioesterase	10.88	R	C1, C3, C4	0.000
VP_011603462.1	FRAAL2298	folD	Methylenetetrahydrofolate dehydrogenase FolD	10.86	Н	А	0.001
VP_011605870.1	FRAAL4755		Membrane phosphatase	10.80	Κ	А	0.000
VP_011607913.1	FRAAL6878	rsmG	Ribosomal RNA small subunit methyltransferase G	10.77	М	А	0.001
VP_011607828.1	FRAAL6795	qorB	Quinone oxydoreductase QorB	10.67	М	C1, C3, C4	0.001
WP_011606846.1	FRAAL5774	dxr	1-Deoxy-D-xylulose 5-phosphate reductoisomerase Dxr	10.54	Ι	А	0.001
VP_050997128.1	FRAAL3342	ssuA	Sulfonate transport system substrate-binding SsuA	10.50	Р	C1, C2, C4	0.000
VP_041938619.1	FRAAL0128	purF	Amidophosphoribosyltransferase PurF	9.86	F	А	0.000
WP_009742300.1			DNA-binding protein HU-beta, NS1 (HU-1)	9.76	L	А	0.016
VP_041939818.1			ATP-binding protein	9.60	-	А	0.004
VP_041939008.1	FRAAL1862	rpiB	Ribose 5-phosphate isomerase RplB	9.07	G	А	0.003
VP_011601953.1	FRAAL0711		Transcriptional regulator	8.76	Κ	C1, C1c, C3, C4	0.002
WP_009737569.1	FRAAL4338	bfrA	Bacterioferritin BfrA Putative	8.49	Р	А	0.004
WP_011602931.1	FRAAL1747		carboxyvinyl–carboxyphosphonate phosphorylmutase	8.43	G	C1, C1c, C4	0.000
WP_041938916.1	FRAAL1429	hpnD	Phytoene synthase HpnD	8.40	Ι	А	0.003
WP_011604375.1	FRAAL3226	bkdC	Branched-chain alpha-keto acid dehydrogenase subunit BkdC	8.14	С	А	0.000
VP_041940119.1	FRAAL0708		Lipid esterase	7.91	R	C1, C2, C3, C4	0.000
WP_011602626.1	FRAAL1431	fppS	Farnesyl diphosphate synthase	7.75	Н	A	0.002
WP_041939587.1			LuxR family transcriptional regulator	7.67	Κ	C1, C1c, C3, C4	0.000
WP_011606077.1		adk	Adenylate kinase	7.50	F	А	0.001
WP_011604508.1			Short-chain fatty acid-CoA ligase	7.43	Ι	А	0.000
WP_011604193.1			Transcriptional regulator	7.36	K	f	0.000
WP_041939680.1	FRAAL5348	cspA	Cold shock protein	7.33	K	A	0.002
WP_011602011.1	FRAAL0771	dgc	Diguanylate cyclase	7.17	Т	f	0.000

Table 1. Com.	Tab	le 1	. Cont	
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NCBI Accession	FRAAL #	Gene	Functional Description	Fold Change	COG	Distribution	% NSAF Nodule
WP_011607242.1	FRAAL6192		Adenylate kinase	7.17	F	А	0.001
WP_041939231.1	FRAAL2835	adh	Acyl-CoA dehydrogenase	6.71	Ι	А	0.000
WP_011606204.1	FRAAL5096	pks	Polyketide cyclase/dehydrase and lipid transport	6.56	-	А	0.001
WP_050997247.1	FRAAL5736	ftsK	Cell division protein FtsK	6.43	D	А	0.000
WP_011604554.1	FRAAL3408		Alpha/beta hydrolase	6.37	R	f	0.001
WP_041938863.1	FRAAL1156	sucC	Succinyl-CoA synthetase subunit beta SucC	6.14	С	А	0.012
WP_011605559.1	FRAAL4438		Sugar-phosphate dehydrogenase	6.08	С	А	0.002
WP_041939760.1	FRAAL5796	smc	Chromosome segregation protein Smc	6.04	D	А	0.000
WP_041939603.1	FRAAL4991	thiF	Thiamine biosynthesis protein ThiF	6.00	Н	f	0.000
WP_041938832.1	FRAAL0997	hemL	Glutamate-1-semialdehyde aminotransferase HemL	5.93	Н	А	0.001
WP_011603632.1	FRAAL2475	glbN	Truncated hemoglobin GlbN	5.87	R	f	0.001
WP_011601393.1	FRAAL0129	purM	Phosphoribosylaminoimidazole synthetase PurM	5.77	F	А	0.001
WP_041938655.1	FRAAL0282		Putative integrase/resolvase	5.71	L	f	0.000
WP_011603454.1	FRAAL2289		Putative two-component oxygen sensor kinase	5.67	Т	А	0.000
WP_011604752.1	FRAAL3613		Diacylglycerol O-acyltransferase	5.67	-	f	0.000
WP_011602534.1	FRAAL1337	pkn	Serine/threonine protein kinase	5.50	Т	F	0.000
WP_011603231.1	FRAAL2064	acnA	Aconitate hydratase AcnA	5.44	С	А	0.006
WP_011605678.1	FRAAL4561	sufE	(2Fe-2S)-Binding protein SufE	5.40	Р	А	0.001
WP_050997107.1	FRAAL2825		Putative N-fatty-acyl-amino acid synthase	5.38	Е	А	0.001
WP_011606156.1	FRAAL5045	ahpE	Peroxiredoxin AhpE	5.31	0	А	0.009
WP_041939398.1	FRAAL3827	bra	Branched-chain amino acid ABC transporter ATP-binding protein	5.18	Е	А	0.001
WP_011607046.1	FRAAL5991	rshA	Anti-sigma factor RshA	5.12	Т	А	0.004
WP_011602689.1	FRAAL1499	cysO	Sulfur carrier protein CysO	5.07	Н	А	0.002
WP_011602425.1	FRAAL1224		Acetyl-CoA synthetase	5.04	Ι	А	0.001
WP_011602282.1	FRAAL1055	pyrE	Orotate phosphoribosyltransferase pyrE	5.00	F	А	0.001
WP_041939967.1	FRAAL6738	purS	Phosphoribosylformylglycinamidine synthase PurS	4.92	F	f	0.001
WP_041939945.1	FRAAL6664	purE	Phosphoribosylaminoimidazole carboxylase PurE	4.90	F	А	0.003
WP_035923438.1	FRAAL6556	crp	Crp/Fnr family transcriptional regulator	4.83	Т	А	0.005
WP_011606876.1	FRAAL5804	rnc	Ribonuclease III rnc	4.82	Κ	А	0.000
WP_011604517.1	FRAAL3370	fklB	Peptidyl–prolyl cis–trans isomerase FklB	4.80	0	А	0.003
WP_041939534.1	FRAAL4560	sufC	Fe-S cluster assembly ATPase SufC	4.62	0	А	0.004
WP_011603306.1	FRAAL2141	ruvB	Holliday junction DNA helicase RuvB	4.60	L	А	0.000
WP_011605202.1	FRAAL4072	pks	Polyketide synthase	4.56	Q	f	0.000
WP_011601665.1	FRAAL0409		Acetyl-CoA acetyltransferase	4.53	Ι	А	0.000
WP_011606938.1	FRAAL5877	etfA	Electron transfer flavoprotein EtfA	4.48	С	А	0.007
WP_011601417.1	FRAAL0156	gdhB	Glutamate dehydrogenase GdhB	4.47	Е	f	0.000

The most underabundant was a cAMP-binding membrane-bound transcriptional regulator (FRAAL6506) at an FC of 0.01 (Supplementary Table S1) in synton (set of genes with a conserved order) with a geosmin synthase gene. There were 2983 proteins detected that belonged to *Frankia alni*. There were also 68 overabundant proteins in nodules and 30 overabundant ones in pure cultures that did not meet the *p*-value criterion (\leq 0.05). There were 729 proteins that were not certified by mass spectrometry with the validation of at least two distinct peptides in this dataset. The overabundant protein coding genes were scattered around the genome (Figure 1).

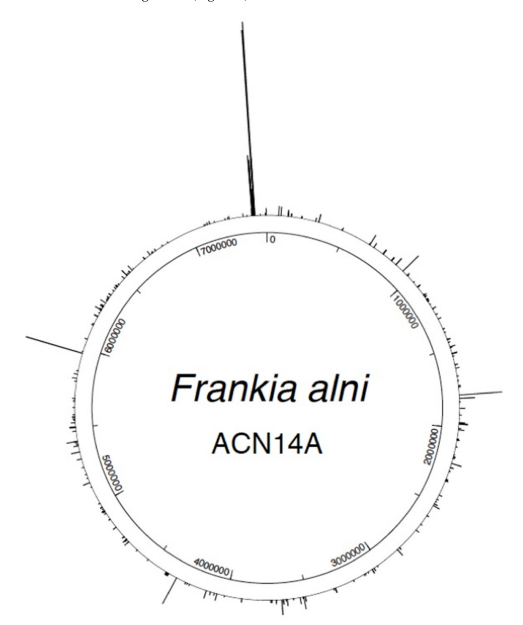


Figure 1. Circular map of the genome of *Frankia alni* with the proteins over-abundant in nodules relative to a nitrogen-replete pure-culture (FC \geq 2) positioned along the genome. The *nif* genes are at the top close to the origin of replication.

Among *F. alni* proteins, the nitrogenase proteins were the most overabundant with 7 among the 10 highest using as reference a nitrogen-replete pure culture. In addition to NifH, NifK (FRAAL6811) at an FC of 279 and NifD (FRAAL6812) at an FC of 86.17, the *nif* cluster comprises 17 genes from FRAAL6798 (*korB*) to FRAAL6814 (*nifV*), 13 of which were identified accounting for 3% of the cumulated NSAF in nodules.

Proteins associated to symbiotic life such as the Suf cluster (FRAAL4558-FRAAL4564, FC = 5.4 - 3.39) and the squalene lipids cluster (FRAAL1427-FRAAL1435, FC = 8.4) were among those detected as overproduced; however, the Hup clusters were not detected (FRAAL1822-FRAAL1832; FRAAL2388-FRAAL2402). Proteins involved in the upstream non-mevalonate pathway for hopanoids biosynthesis such as Dxs (FRAAL2088, FC = 2.34), Dxr (FRAAL5774, FC = 10.54), IdsA (FRAAL1431, FC = 7.75) and Idi (FRAAL6504, FC = 2.21) were also overproduced.

Other lipid metabolism proteins were overabundant such as several unlinked acetyl-CoA acetyl transferases (FRAAL4934, FC = 12.91; FRAAL2618, FC = 11.83; FRAAL0409; FC = 4.53; 3973, FC = 2.95), a fatty acid-CoA ligase (FRAAL3361, FC = 7.43), an acyl-CoA dehydrogenase (FRAAL2835, FC = 6.71) and a diacylglycerol O-transferase (FRAAL3613, FC = 5.67).

Oxidative stress proteins such as SodF (FRAAL4337, FC = 3.85) and its bacterioferritin neighbor (FRAAL4338, FC = 8.49), glutathione peroxidase (FRAAL1783, FC = 3.55), two nitroreductases (FRAAL4025, FC = 44.33; FRAAL3236, FC = 18.83) and rubrerythrin (FRAAL5116, FC = 3.08) were overabundant. Other stress proteins were underabundant such as KatA (FRAAL3167, FC = 0.09) and glutathione synthase (FRAAL4656, FC = 0.03) or were not identified such as KatB (FRAAL3889).

Several TCA cycle proteins such as succinyl-CoA synthase (FRAAL1156, FC = 6.14), aconitate hydratase (FRAAL2064, FC = 5.44), malate dehydrogenase (FRAAL5612, FC = 4.37) or fumarase (FRAAL6152, FC = 3.96) were more abundant in nodules, but several others were not detected. There were 15 ribosomal proteins overabundant at an FC \geq 2 (FRAAL1069, FRAAL1068, FRAAL1097, etc.) as well as other ancillary proteins such as a ribosome-recycling factor (FRAAL5778, FC = 3.35).

Other carbohydrate metabolism proteins were overabundant such as glycosyl transferases (FRAAL2049, FC = 15.5; FRAAL4552, FC = 3.5; FRAAL5400, FC = 2.85), a sugar epimerase (FRAAL6795, FC = 10.67), a ribose isomerase (FRAAL1862, FC = 9.07), a sugarbinding protein (FRAAL5914, FC = 3.49) and an enolase (FRAAL6233, FC = 3.46).

Ammonia assimilation proteins were overabundant such as the nitrogen regulatory protein P-II (FRAAL0904, FC = 3.27), glutamine-synthetase adenylyltransferase (FRAAL5164; FC = 2.13), glutamate-1-semialdehyde aminotransferase FRAAL0997, FC = 5.93) and a GS (FRAAL6426; FC = 1.8), but several others such as GSII were not detected or were underabundant such as GSI (FRAAL5161, FC = 0.01) and GOGAT (FRAAL4964-65, FC = 0.08 - 0.05).

A cell-division control ATPase was overabundant FRAAL5516 (FC = 89), while other proteins with related functions were less abundant (FRAAL6588, cell division protein FtsH FC = 0.22) or not detected.

Chaperones were overabundant such as a disaggregation chaperone (FRAAL2601, FC = 4.22) and several GroEL homologs (FRAAL6640, FC = 2.64; FRAAL1699, FC = 2.16; FRAAL1133, FC = 2.2; FRAAL6486, FC = 1.94). Other overabundant proteins with related functions were FRAAL4269 (Clp Protease), FRAAL6701 (GroEL) and FRAAL5348 (cold shock), with the corresponding genes scattered on the chromosome.

Some transporters were overabundant in nodules such as a molybdenum transporter (FRAAL1673, FC = 64.17), a BCA transporter (FRAAL3827, FC = 5.18) and a C4-dicarboxylate transporter (FRAAL1390, FC = 3.42). A phosphate transporter (FRAAL6535) was also overabundant (FC = 2.09), while many were underabundant.

Some regulators were overabundant in nodules such as a Tet-R (FRAAL0868, FC = 33.09) and a Sarp-R (FRAAL0301, FC = 12.38). A Bhl-R (FRAAL5827, FC = 9.76), a MarR (FRAAL0711, FC = 8.76) and a luxR (FRAAL4898, FC = 7.67) were also overabundant, while many were underabundant. An anti-sigma factor (FRAAL5991, FC = 5.12) in synton with a sporulation-specific sigma factor was overabundant.

Most secondary metabolite synthesis proteins were not detected or underabundant except two PKS (FRAAL5096, FC = 7.22; FRAAL4072, FC = 4.56) and a bacteriocin (FRAAL4503, FC = 2.16).

A gamma-aminobutyraldehyde dehydrogenase (FRAAL6022, FC = 3.73) was overabundant. This enzyme transforms 4-aminobutanal into 4-aminobutanoate (GABA).

The Kdp proteins (FRAAL5299-FRAAL5305), the cellulase synton proteins (FRAAL4954-FRAAL4959) and the acyl-dehydrogenases (FRAAL1659-FRAAL1670) were not identified.

Almost all carboxylases were underabundant such as three acetyl-CoA carboxylases (FRAAL3159, FC = 0.55; FRAAL1213, FC = 0.43; FRAAL3158, FC = 0.23), a methyl-crotonoyl-CoA carboxylase (FRAAL2343, FC = 0.15) and two propionyl-CoA carboxylases (FRAAL5672, FC = 0.05; FRAAL1210, FC = 0.68). The only overabundant one was a phosphoribosylaminoimidazole carboxylase (FRAAL6664, FC = 4.9) involved in the synthesis of purines.

A COG analysis (Figure 2) revealed that "C" (energy production) and "J" (translation) were the most represented in the nodule overabundant proteins, while "E" (amino acid transport and metabolism) and "K" (transcription) were the most represented in the underabundant proteins (Figure 2). Many COGs were very rare in the nodule (N, cell motility; U, intracellular trafficking; V, defense mechanisms and Q, secondary metabolites).

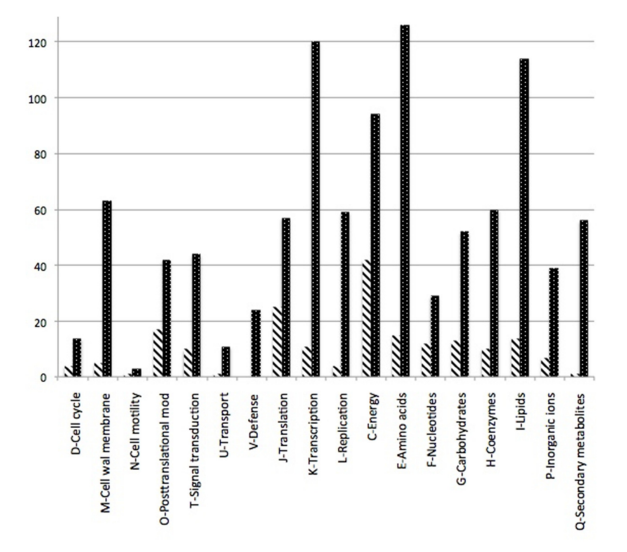


Figure 2. Histogram of over- and underabundant proteins coding genes (FC > 2 hatched; FC < 0.5 dotted) assigned to COGs. Undefined categories and those with one or less occurrences were omitted.

A drawing with the major overabundant proteins in nodules permits to show nitrogenase and other energy-generating, transporter and stress-coping determinants (Figure 3).

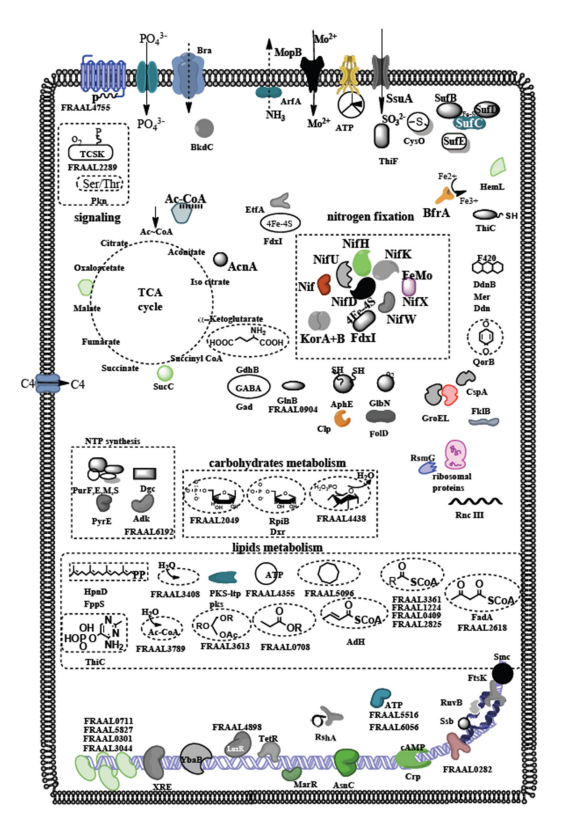


Figure 3. Schematics of a *Frankia* cell in its symbiotic host cell showing the likely function and localization of the 100 most abundant proteins related to nitrogen fixation, lipid metabolism, Krebs cycle (TCA), cell signaling, electron transfer, transport of anions and cations, enzymes involved in nucleotides biosynthesis, DNA metabolism and regulation and storage and stress proteins. The identities of the proteins and specific genes are listed in Table 1.

4. Discussion

Proteomics has the potential to decipher the physiological changes occurring in cells upon ecological transitions [32]. Establishment of the actinorhizal symbiosis is a drastic change for both partners that cannot be analyzed through negative genetics. Omics thus offer an interesting approach to pinpoint the key molecular players from both organisms. Proteomics in particular has been used to study *Frankia* either in early interaction with *Alnus* [18]—or with *Elaeagnus angustifolia, Ceanothus thyrsiflorus* and *Coriaria myrtifolia* [17], showing, among others, cellulose synthase and a potassium transporter over-detected upon contact with *Alnus* [18] and nitrogen fixation and assimilation proteins upon contact with *Elaeagnus* [17], as well as stress response and respiration proteins—or in the mature field nodules of a range of actinorhizal plants [21].

Nitrogenase is the most abundant protein complex in symbiotic *Frankia alni*. It comprises 17 genes that had previously been shown to be the most transcribed in mature nodules [7]. The present study confirms our vision that symbiotic *Frankia* is essentially a nitrogenase machine. Associated proteins were seen here, such as the SuF cluster assembly proteins and the hopanoid biosynthesis proteins (that protect nitrogenase from oxygen), but not the hydrogenase proteins even though they were among those highly expressed [33]. *Frankia* hydrogenase proteins seen previously in field nodules [21] were not detected.

The high overabundance for 2-oxoglutarate ferredoxin oxidoreductase (KorAB) confirms a role initially hypothesized based on its position next to *nif* genes of 2-oxoglutarate as a primary electron source for nitrogenase [21]. The position of the *korAB* genes next to *nif* genes and their duplication in symbiotic strains only are probably the result of a specialization on the one hand for symbiosis (FRAAL6798-6790, FC = 91.5) and on the other hand for saprophytic life (FRAAL1050-1051, FC = 0.80) as seen before for Hup [34].

Nitrogen fixation is an energy-demanding process that consumes eight ATP molecules per molecule of NH_4^+ produced. It is, thus, expected that the TCA cycle will be running at full force, transforming the photosynthates given by the plant into energy. The transcriptome of symbiotic *F. alni* was seen to contain several TCA genes upregulated at a high level [7] as was the case for the transcriptome of *F. soli* upon early contact [17].

There is an important specialization in nodules with only a twentieth of the genome identified (250 proteins with an FC \geq 2) relative to a pure culture where five times as many proteins were identified at the same threshold. A similar specialization was noted in *S. meliloti* nodules [35].

The oxidative stress proteins catalase and superoxide dismutase are essential for the nodulation of *Sinorhizobium* [36] to cope with the oxidative burst induced as an early plant defense response against avirulent pathogens [37]. Nitroreductases play a similar role in *Sinorhizobium* [38]. In actinorhizal nodules, there is also an oxidative burst with which *Frankia* must cope [39]. Pure cultures of *F. alni* have been shown to have a basic expression of its two catalases that are then markedly upregulated upon contact with H_2O_2 or with methyl viologen [40]. However, it appears that this upregulation is not maintained in mature nodules.

GABA is considered a nutrient as well as an effector to trigger plant responses to heat, salt, herbivory and other stresses [41]. GABA has been detected in the metabolome of *Alnus* and *Casuarina* nodules and roots at very high levels [42,43]. GABA was also seen to improve nitrogenase and respiration in pure culture [43]. In leguminous nodules, GABA has been suggested to be a nutrient fed to the symbiont since it accumulated labeled $^{15}N_2$ and contributed to the N-nutrition [44]. In *Alnus*, labeled $^{15}NH_4^+$ was recovered as alanine (Ala), γ -amino butyrate GABA), glutamine (Gln), glutamate (Glu), citrulline (Cit) and arginine (Arg) [45]. GABA was late to appear, but then, it continued at a fast pace. Since GABA synthesis involves the utilization of protons and releases CO₂, it has been suggested as a means to reduce acidity [46].

The photosynthates fed by the plant to the bacterium have been the subject of several hypotheses. It has long been known that *Alnus*-infective strains could not use sugars [3], which is why organic acids are routinely used to grow most strains. The identification

of a dicarboxylate transporter among those genes specific to nodules [47] indicated that dicarboxylates were good candidates. Several of these were measured in nodules and roots, and fumarate, succinate, malate and 2-oxoglutarate were found to be overabundant in nodules, although this approach is biased by the presence of mitochondria where citrate is also very abundant, yet it is toxic to *Frankia* [48]. The overabundance in the present study of the C4-dicarboxylate transporter, FRAAL1390 confirms that dicarboxylates play a role in trophic exchanges between symbionts, although its modest overabundance (FC = 3.42) points either to other transporters or to other photosynthates involved.

Chaperones such as GroEL have been shown to be essential to nodulation in *Sinorhizo-bium meliloti* [49]. They are also involved in response to other stresses such as heat [50] or salt [51] or simply for the assembly of very complex proteins such as nitrogenase [52]. Such chaperones were previously identified in the transcriptome of symbiotic *Frankia alni* [7].

Proteomics permits complementation of the vision achieved by transcriptomics. The two approaches have different strengths and weaknesses. Shotgun proteomics carried out in the standard mode (i.e., based on trypsin proteolysis) may miss small proteins, membrane proteins and polypeptides with rare combinations of arginine and lysine residues [53]. Transcriptomics misses unstable messengers or those with secondary structures. Furthermore, proteomics and transcriptomics address molecular entities with different half-lives and dynamics, thus they are highly complementary. The present study reinforces the picture of a microbe highly specialized for the energy-demanding nitrogenase and the accompanying stresses. It sheds no light, however, on the molecular processes involved in communication between the two partners.

Supplementary Materials: The following supporting information can be downloaded at https: //www.mdpi.com/article/10.3390/microorganisms10030651/s1: Table S1: List of *Frankia* proteins identified in nodules and in a nitrogen-replete pure culture and their spectral counts.

Author Contributions: P.P., N.A. and P.N. conceived the experiment; P.P., N.A. and P.F. grew the plant and the bacterium; G.M. and J.A. extracted the proteins and ran the proteomic analysis; D.A. and P.N. did the analyses and P.P. and P.N. wrote the paper. All authors have read and agreed to the published version of the manuscript.

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