Alfalfa snakin-1 prevents fungal colonization and probably coevolved with rhizobia

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Abstract

Background: The production of antimicrobial peptides is a common defense strategy of living cells against a wide range of pathogens. Plant snakin peptides inhibit bacterial and fungal growth at extremely low concentrations. However, little is known of their molecular and ecological characteristics, including origin, evolutionary equivalence, specific functions and activity against beneficial microbes. The aim of this study was to identify and characterize snakin-1 from alfalfa (MsSN1).

Results: Phylogenetic analysis showed complete congruence between snakin-1 and plant trees. The antimicrobial activity of MsSN1 against bacterial and fungal pathogens of alfalfa was demonstrated in vitro and in vivo. Transgenic alfalfa overexpressing MsSN1 showed increased antimicrobial activity against virulent fungal strains. However, MsSN1 did not affect nitrogen-fixing bacterial strains only when these had an alfalfa origin.

Conclusions: The results reported here suggest that snakin peptides have important and ancestral roles in land plant innate immunity. Our data indicate a coevolutionary process, in which alfalfa exerts a selection pressure for resistance to MsSN1 on rhizobial bacteria. The increased antimicrobial activity against virulent fungal strains without altering the nitrogen-fixing symbiosis observed in MsSN1-overexpressing alfalfa transgenic plants opens the way to the production of effective legume transgenic cultivars for biotic stress resistance.

Keywords: Antimicrobial peptides, Alfalfa, Evolution, Land plants, Innate immunity, Snakin

Background

Alfalfa (Medicago sativa L.), known as the “Queen of Forages”, is a perennial legume. This species is native to Asia, and is considered one of the first known crops with a cultivation history of at least 3500 years. Due to its strong vitality, high nutritional quality, high yields, high adaptability and multiple uses, alfalfa is the main forage crop produced in temperate regions of the planet. Elite alfalfa cultivars must not only have high forage yields but also maintain their productivity and stands over several years to provide substantial economic benefits. Regarding this complex topic, improved fungal disease resistance has been specifically identified as the critical trait in alfalfa persistence [1]. In this context, it is proposed that the use of snakin-1 peptide (SN1), a powerful but poorly studied antimicrobial compound, to improve alfalfa tolerance to virulent fungal pathogens should be explored.

Antimicrobial peptides are present in virtually all organisms and are an ancient and critical component of innate immunity. SN1, the first member of the snakin family to be characterized, was isolated from a crude cell wall preparation of potato (Solanum tuberosum) tubers (StSN1) [2]. This cysteine-rich peptide from potato was found to be active against bacterial and fungal pathogens at extremely low concentrations (EC50 < 10 μM) [3]. The expression pattern of the StSN1 gene suggests that plant SN1 could be a component of constitutive defense barriers, especially those of storage and reproductive plant organs [3]. A second snakin peptide (StSN2), which has a high amino acid identity (30%) to StSN1, was also isolated from a crude cell wall preparation of potato tubers. Consistent with this high amino acid identity, StSN2 is also active at very low concentrations against a wide range of pathogens [4]. In contrast to StSN1, the expression
of StSN2 is locally induced by wounding and pathogen infection, suggesting a critical role of snakin-2 in both constitutive and inducible defense barriers of plants. These strong antimicrobial activities of snakin peptides have been verified using bacterial and eukaryotic heterologous expression systems [5-7]. Based on the presence of the Gibberellic Acid Stimulated Arabidopsis (GASA) domain and the absence of bioinformatic (e.g. RGD residues) and functional (e.g. toxic activity) data supporting its relationship with cysteine-rich peptides from snakes venoms, StSN1 and StSN2 have been recently renamed as GSL1 and GSL2, respectively [8,9].

Overexpression of SN1 in potato and wheat (Triticum aestivum), SN2 in potato and tomato (Solanum lycopersicum), and snakin-defensin hybrid protein in tobacco (Nicotiana tabacum) and potato restricts pathogen invasiveness and enhances tolerance to bacterial and fungal diseases, without altering the agronomic phenotype of these crops [6,9-12]. Furthermore, disease sensitivity is enhanced by silencing SN2 in wild tobacco (Nicotiana benthamiana), supporting the central role of snakin peptides in plant defense [13]. In addition, it has been recently shown that StSN1 is located in the plant cell wall [14], confirming that snakin peptides are components of the physical barrier and the first line of defense used by plant cells to prevent bacterial and fungal entry [3]. Furthermore, in concordance with the functional classification of snakin peptides as members of the GASA protein family [4,15,16], StSN1 silencing affects cell division, primary metabolism, and cell wall composition [14]. Snakin peptides could have additional functions in plant growth and development beyond their demonstrated function in biotic stress response. In spite of their hypothetical functional similarity, there is little to no phylogenetic reports of the relationship between StSN1 and GASA-related proteins.

The aim of this study was to identify and characterize snakin-1 (MsSN1) gene of alfalfa. The phylogenetic and functional analyses showed here, propose that MsSN1 is an ancestral plant gene involved in biotic stress resistance, suggesting a coevolutionary process, in which alfalfa exerts a selection pressure for resistance to MsSN1 on rhizobial bacteria.

**Methods**

**Bacterial and fungal strains**

The bacterial strains (all Gram-negative bacteria) used in this study were: *Pseudomonas fluorescens* Pf-5 [17], *Sinorhizobium meliloti* BL225C [18], *Sinorhizobium meliloti* SM11 [19], *Sinorhizobium medicae* WSM419 [20], *Sinorhizobium fredii* USDA 257 [21], *Rhizobium* sp. Or 191 [22], *Rhizobium etli* CFN 42 [23], *Mesorhizobium loti* MAFF303099 [24], *Bradyrhizobium japonicum* USDA110 [25] and *Agrobacterium tumefaciens* LBA4404 [26]. The fungal strains used in this work were: *Phoma medicaginis* strain CT1 and *Colletotrichum trifolii* strain CT2, isolated from INTA alfalfa cultivars and kindly provided by Dr. Ricardo Comerio (Instituto de Microbiología y Zoología Agrícola, Instituto Nacional de Tecnología Agropecuaria, Argentina).

**Fungus material**

Two fungal pathogens, *Phoma medicaginis* strain CT1 and *Colletotrichum trifolii* strain CT2, were grown on PDA (Cat. # B0216605, Britania) plates at room temperature for approximately 7 days before the start of the bioassay. For spore collection, the plates were flooded with sterile distilled water and scraped with a wire loop. Spore concentration was adjusted to $1 \times 10^6$ spores/ml for *C. trifolii* CT2 and to $1 \times 10^5$ spores/ml for *P. medicaginis* CT1 with sterile distilled water. The fungal strains were maintained through sequential passages in plants.

**Plant material**

The *Medicago sativa* plants used were the regenerative clone C2-3, kindly provided by Drs. B. McKersie and S. Bowley (Plant Biotechnology Division, Department of Plant Agriculture, University of Guelph, Canada), and the regenerative clone 432-19-17, previously isolated in our laboratory.

**Bacterial and plant RNA Isolation and cDNA synthesis**

Total bacterial (*E. coli*) and plant tissues (roots, stem, leaflets) RNA was extracted by using an RNeasy Mini Kit (Cat. # 74106, Qiagen) following the manufacturers’ instructions. Samples of 2 μg total RNA isolated from bacterial cells or plant tissues were reverse-transcribed in a 25 μl reaction using MMLV-RT (Cat. # M1701, Promega). For PCR amplification in bacteria and plants, 1 μl of RT reaction was used. The PCR reactions were carried out in 25 μl with 0.5 μM of each primer [27], using Taq polymerase (Cat.# 11615010, Invitrogen) following the manufacturers’ instructions.

**MsSN1 plasmid construction**

The MsSN1 cDNA (GenBank accession number JQ809686) was isolated using primers p1 FW and p2 RV (Additional file 1) designed against the 5′ and 3′ untranslated regions (UTR) of the putative SN1 gene (GenBank accession number XM_003589066, MTR_1g018640) from *Medicago truncatula*. Full-length cDNA was amplified by PCR and this fragment was cloned into a pCR2.1TOPO vector (Cat. # K4500-01, Invitrogen). The pTOPO-MsSN1 plasmid was digested with Not I (Cat. # R6431, Promega), treated with Klenow, and religated to destroy the polylinker Not I site. The resulting plasmid was named pTOPO-NotI-MsSN1. The sequencing reactions of pTOPO-NotI-MsSN1 were performed at INTA-Argentina (www.inta.gov.ar).
The cDNA sequence was named MsSN1 (*Medicago sativa* snakin-1). To produce recombinant bacteria expressing MsSN1, plasmid pSJ33-MsSN1 carrying MsSN1 was constructed by subcloning the 0.5-kb EcoRI (Cat. # R6011, Promega) fragment from pCR2.1TOPO-MsSN1 into pSJ33 [28], and afterward introduced in *Escherichia coli* for heterologous expression of MsSN1. To produce transgenic alfalfa lines overexpressing MsSN1, pCR2.1TOPO-Not1-MsSN1 was digested with KpnI (Cat. # R6341, Promega) and XbaI (Cat. # R6181, Promega), and the MsSN1 restriction fragment was cloned into pKANNIBAL vector (GenBank accession number AJ311873). The resulting plasmid MsSN1 was digested with NotI, and the 3SSMsSN1 restriction fragment was cloned into pART27 binary vector [29]. The resulting recombinant binary vector containing the MsSN1 cDNA with its signal peptide under the CaMV 35S promoter was named pART-35S::MsSN1.

**Bioinformatic analysis of MsSN1**

MsSN1 sequence of *Medicago sativa* (AFE82743), a peptide composed of 91 amino acids, was used as query to search against all available complete eukaryotic genome databases in NCBI with protein annotation in GenBank. The cut-off to obtain candidate orthologs was 20% of amino acid identity (Additional file 2). Sequences were searched using by BLASTP tools in NCBI and PLAZA databases (http://www.ncbi.nlm.nih.gov/blast; http://bioinformatics.psb.ugent.be/plaza). Protein identity calculations were performed using MatGAT v2.02 [30]. Evolutionary analysis was conducted by using MEGA version 5.0 [31]. Protein sequences were aligned using the ClustalW program. Phylogenetic trees were constructed using the neighbor-joining method with genetic distances computed using the pairwise deletion model and bootstrap analysis of 500 values and root on midpoint (i.e. midpoint of the longest pathway between two clusters of sequences). *In silico* analysis of conserved motives in putative snakin/GASA proteins, Pfam domains and signal peptides were predicted by using Pfam and Signal-3 L with default parameters, respectively [32,33].

**DNA extraction and sequence analysis of fungal strains**

Fungal DNA was extracted as previously described by Moller [34] with modifications suggested by Dr. E.W. Boehm (http://www.ebohm.com/). Briefly, 100 mg of fungal mycelia was scraped from 10-day-old PDA cultures, ground in a 1.5 ml tube with micropestle by adding 500 μl of Lysis Buffer (100 mM Tris pH 8, 10 mM EDTA, 2% SDS, 1% β-Mercaptoethanol, 100 μg/ml proteinase K). Lysate was incubated at 60°C for 60 min. Subsequently, 5 M NaCl was added to a final concentration of 1.4 M and mixed before adding 0.1 vol of CTAB 10% (w/v). The mixture was incubated at 65°C for 10 min. DNA was extracted by adding an equal volume of chloroform:isoamyl alcohol (24:1), incubating for 30 min at 0°C and centrifuging at 14,000 × g for 10 min at 4°C. The aqueous phase was mixed with 0.5 vol of 5 M ammonium acetate, incubated at 0°C for 60 min and centrifuged at 14,000 × g for 1 min. DNA was precipitated from the supernatant by adding 0.55 vol of isopropanol, centrifuged at 14,000 × g for 10 min and washed twice with 70% ethanol. DNA pellet was air-dried and resuspended in 50 μl of TE (10 mM Tris pH 8.0; 1 mM EDTA pH 8.0). The internal transcribed spacer regions 1 & 2 and 5.8S nrDNA (ITS) and partial sequences of TUB genes were amplified and sequenced by Macrogen Inc. (Korea) service (http://www.macrogen.com/) using the primer pairs ITS-1FW and ITS-4RV and Btub2FW and Btub4RV, respectively (Additional file 1). The nucleotide sequences obtained here were deposited in the EMBL Nucleotide Sequence Database, accession numbers: KF846005 for *C. trifolii* TUB, KF846006 for *P. medicaginis* TUB, KF846009 for *C. trifolii* ITS and KF846010 for *P. medicaginis* ITS.

**In vitro antimicrobial activity assays**

*Escherichia coli* recombinant strains containing the pSJ33 empty vector or pSJ33-MsSN1 were grown overnight at 30°C with shaking (250 rpm) in LB medium supplemented with 1 mM isopropyl-β-D-thiogalactopyranoside in 250-ml Erlenmeyer flasks containing 50 ml of medium. Samples of 25 ml of culture broth (O.D: 0.8) were centrifuged at 5,000 × g at 4°C for 10 min. The pellets were washed three times with physiological solution (0.9% NaCl). Crude cell lysates were achieved by four consecutive cycles of freezing in liquid nitrogen followed by thawing at 37°C. After centrifugation at 14,000 × g at 4°C for 20 min, the supernatant was resuspended in 400 μl of physiological solution. For the analysis of MsSN1 expression in *E. coli*, total bacterial RNA was extracted by using the RNAeasy Mini kit and treated with DNasel (Cat. # M6101, Promega). cDNA was obtained using random hexamers (Cat. # B070-40, Promega) and AMV Reverse Transcriptase (Cat. # M9004, Promega). For PCR amplification, 1 μl of RT reaction was used. The PCR reactions were carried out in 25 μl with 0.5 μM of each primer according to Setten [27]. PCR conditions comprised: 1 cycle at 94°C for 3 min, 34 cycles of 94°C for 45 s, 56°C for 1 min and 72°C for 1 min. The expression of MsSN1 was analyzed using primers p3 FW and p4 RV which amplify the complete open reading frame (Additional file 1).

The disk inhibition assays were evaluated as described by Ayub [35], with very slight modifications. Cultures were performed in 125-ml Erlenmeyer flasks containing 25 ml of TY medium [36] or LB medium for rhizobia or *Pseudomonas* and *Agrobacterium*, respectively. Bacteria were incubated overnight at 28°C with shaking (250 rpm). Sterile Whatman No. 1 filter disks (5 mm) impregnated with 5 μl of MsSN1-free (*E. coli* pSJ33) or MsSN1 (*E. coli* pSJ33) were cut in a 1.5 cm diameter and loaded into the wells of a 96-well microplate. The wells were filled to a volume of 200 μl with sterile LB medium for rhizobia and TY medium for *Pseudomonas* and *Agrobacterium*.
μ fresh medium every 2 weeks, at 25°C and 16 h light (100 μM acetosyringone for 2 days at 25°C in the dark).

Plant transformation
The recombinant binary vector pART-35S::MsSN1 was introduced into Agrobacterium tumefaciens LBA 4404 by electroporation, by using the procedure described by Shen & Forde [37]. Petioles of alfalfa clone C2-3 were transformed with pART-35S::MsSN1 via A. tumefaciens and cultured in vitro as described by D’Halluin [38], with slight modifications (Additional file 3). Petiole tissues were decontaminated by immersing in 70% ethanol for 1 min and then in 2% sodium hypochlorite for 20 min. The petioles were washed 3 times in sterile distilled water. Explants previously injured with a scalpel were inoculated with a bacterial culture for 2 minutes (OD₆₀₀ mm = 0.5-0.8), and then dried in Whatman filter paper and transferred onto a solid co-cultivation SHK medium [3% sucrose, 0.435% KSO₄, 2 mg/l 2,4-D, 0.2 mg/l kinetin, 6.5 g/l agar, 20% SHK stock solution (w/v) pH:5.8 (300 mg/l NH₄H₂PO₄, 2.5% KNO₃, 200 mg/l CaCl₂·2H₂O, 400 mg/l MgSO₄·7H₂O, 4.3% K₂SO₄, 1 mg/l KI, 5 mg/l H₂BO₃, 10 mg/l MnSO₄·H₂O, 1 mg/l ZnSO₄·H₂O, 1 mg/l Na₂MoO₄·2H₂O, 1 mg/l CuSO₄·5H₂O, 0.1 mg/l CoCl₂, 6H₂O, 26.29 mg/l NaFeEDTA·H₂O, 288 mg/l proline, 53 mg/l thiprolidine, 200 mg/l myo-inositol, 5 mg/l nicotinic acid, 0.5 mg/l pyridoxine, 5 mg/l thiamine)] containing 100 μM acetosyringone for 2 days at 25°C in the dark. The explants were then washed with 0.5 g/l cefotaxime supplemented with distilled water and transferred to selection/induction medium SHK containing 25 mg/l kanamycin and 400 mg/l cefotaxime with routine transfers to fresh medium every 2 weeks, at 25°C and 16 h light (100 μmol·m⁻²·s⁻¹). Somatic embryos were obtained three months later and then transferred to MS rooting medium, composed of Murashige and Skoog Basal Medium (Cat. # M5519, Sigma) diluted 1:2 with water. Transgenic plants were obtained about 6 months after callus induction. The regenerated plantlets were transferred to the greenhouse once they were well rooted. All plants were grown in a greenhouse at 25-20°C day/night temperature and watered daily. Alfalfa transgenic events (named S1, S2 and S3) were obtained 6 months later and then transferred to the greenhouse at 25-20°C day/night temperature and watered daily. All plants were grown in a greenhouse at 25-20°C day/night temperature and watered daily. Alfalfa transgenic events (named S1, S2 and S3) were propagated by crossing with clone 432-19-17 and by cuttings to increase the number of plants available for biochemical, physiological, and genetic analysis.

Identification of transgenic plants
DNA was isolated from leaf tissue with the DNeasy plant mini kit (Cat. # 69104, Qiagen). Transgenic plants were first identified by PCR with primers p5 FW and p6 RV (Additional file 1), designed against the 35S promoter and SN1 regions of recombinant binary vector. The expression of the transgene was corroborated by RT-PCR, using primers p7 FW and p4 RV, designed against the pCR2-1-TOPO-MCS-derived 5′ UTR region and the 5′ UTR of the MsSN1 gene (Additional file 1). All PCR reactions were performed with Taq (Cat. # 11615010, Invitrogen).

For Southern hybridization analysis, genomic DNA was isolated from leaf tissues of greenhouse-grown plants using the DNeasy Plant Maxi kit (Cat. # 68161, Qiagen) following the manufacturer’s indications. DNA was digested with the KpnI restriction enzyme, which cleaves the construct only once. Then, 20 μg of DNA from each sample was digested overnight and blotted after separation on 1% (w/v) agarose gel 1× TAE. The DNA fragments in gels were transferred to a positively charged Nylon membrane (Cat. # 1120972001, Roche). Nylon membranes were crosslinked and then used for hybridization with a DIG-labeled probe. Prehybridization and hybridization was carried out according to the manufacturer’s instructions. The hybridization probe MsSN1-digoxigenin-labeled DNA was generated by PCR by using the PCR DIG probe synthesis kit (Cat. # 11573152910, Roche), using primers p8 FW and p9 RV and then used as a probe (Additional file 1). PCR amplification was performed under standard conditions (25 μl volume using 0.8 μM of each primer, 1X PCR buffer, 0.2 mM each dNTP, 2 mM MgCl₂ and 20 ng of template) with a program of 34 cycles of 94°C for 1 min, 50°C for 30 s and 72°C for 2.5 min and a final cycle of 72°C for 10 min.

Real-time quantitative RT-PCR (RT-qPCR)
For RT-qPCR, PCR amplification was performed with 5 μl of RT (1:5 diluted per reaction, by using 1 U iQ SYBR green Supermix (Cat. # 170–8880, Bio-Rad) and 0.2 mM primers, with the iCycler iQ system. Primers for the real-time qPCR were p10 FW and p11 RV (Additional file 1). qPCR conditions comprised: 1 cycle at 94°C for 5 min, 34 cycles of 94°C for 45 s, 59.1°C for 30 s, and 72°C for 30 s. At each cycle, accumulation of PCR products was detected. The amplification fragment was sequenced and found to be identical (100% nucleotide identity) to the MsSN1 gene. The expression level of MsSN1 was normalized using aspartate aminotransferase (ATT) (AAAB46610) as a housekeeping gene, using primers p12 FW and p13 RV (Additional file 1). The efficiency of primer binding was determined by linear regression by plotting the cycle threshold value versus the log of the cDNA dilution [39]. RT-qPCR experiments were performed two or three times.
with independent RNA samples (biological replicates). For each biological replicate the qPCR reactions were carried out in duplicate.

**Bioassays. In vitro challenges**

Assays were performed in healthy 15 day-old nonfumigated leaflets. Leaflets were decontaminated washing them in flasks with sterile water with Tween-20 (0.01%) for 10 min. After three washes with sterile water, the leaflets were transferred to a 5% hypochlorite solution for 5 min. Finally, leaflets were washed three times with sterile water and transferred to agar-water petri dishes. Plates were maintained in growth chambers programmed for 16 h light at 23°C and 8 h dark at 20°C. Leaflets were inoculated by putting 5 μl of *P. medicae*is CT1 or *C. trifolii* CT2 spore solution per leaflet. Infection evolution was observed and documented by photos and the following damage score was generated to quantify injuries: 1. Healthy leaflet, 2. Countable injuries, 3. Uncountable injuries, 4. Chlorosis, 5. Completely damaged.

**Bioassays. In vivo challenges**

Alfalfa transgenic seeds were obtained by crossing transgenic plants S1, S2 and S3 with 432-19-17 genotype plants. Transgenic and wild type seeds were treated with sulfuric acid for 10 minutes, washed three times with sterile water and placed in petri dishes with 1% agar water at 16 h of light (100 μmoles/m²s) and 25°C. Germinated plantlets were transferred to MS 0.5X flasks and incubated at 25°C with 16 h photoperiod for 1 month, after which plants were transferred to 1:1 vermiculite:perlite containing 12 cysteine residues in conserved positions (SIGMA) to conserve the humidity. Two-month-old plants were inoculated with *P. medicae*is CT1 by spraying spore suspension to all aerial tissues. Percentage of diseased leaflet was analyzed 30-day post-inoculation and the number of plants with regrowth and the percentage of highly defoliated plants was counted 60-days post-inoculation. As an additional severity parameter of plant disease, the percentage of vigor affected plants (i.e. with visual signs of turgidity loss compared with non-inoculated plants) was estimated. All experiments were carried out with at least 10 independent plants per treatment and with at least 5 non inoculated wild type plants used as control.

**Evaluation of bacterial root colonization**

For the symbiosis assay, wild type and transgenic alfalfa seeds were grown in 100% vermiculite and daily irrigated with the minimal medium called “INTA13 without nitrogen” [27]. 10-days-old plants were inoculated with an early stationary-phase culture of rhizobial suspension (*Sinorhizobium melloti* BL225C). After a month, the plants were harvested and the number of pink nodules was analyzed. Non-inoculated plants were used as controls. Three replicates were analyzed for each treatment. The *P. fluorescens* Pf-5 colonization assay was evaluated according to Sanchez [40], with two slight modifications: alfalfa was grown in INTA13 supplemented with Ca(NO₃)₂ [27] and 0.5 NE2 medium plus sodium octanoate (0.25% w/v) was used to estimate the colony forming units [28].

**Statistical analysis**

Biological measurements were repeated at least three times with at least 10 different plants. Significant differences between treatments were calculated using Student’s t-test. qPCR experiments were independently performed three times, with comparable results. The three PCR reactions were carried out in duplicate. Significant differences were calculated using ANOVA followed by Tukey test.

**Results and discussion**

**Identification and evolutionary analysis of MsSN1**

To analyze the evolution and function of MsSN1, a 276-bp cDNA fragment was isolated from roots by RT-PCR, using specific primers designed for a putative snakin-1 peptide predicted from the genome of the model legume *Medicago truncatula* (MsSN1, MTR_1g018640). The PCR product was cloned into a pCR2.1TOPO vector and sequenced (JQ517286). This cDNA fragment shares 98% identity with the putative snakin-1 peptide from *Medicago truncatula* (XP_003589114), hence it was named MsSN1 (*Medicago sativa* snakin-1). Like potato snakin-1 peptide (StSN1), MsSN1 has a putative signal peptide of 25 amino acids and possesses a snakin/GASA domain (Pfam02704) containing 12 cysteine residues in conserved positions within a conserved C-terminal region (Figure 1).

To perform a phylogenetic analysis of SN1, the evolutionary study was restricted to well-characterized sequenced species of plants using proteins with high amino acid identity (>20%). Using this strict criterion, complete congruence (i.e. same topology) between SN1 and vascular plant trees was observed (Figure 2), suggesting that plant SN1 was acquired by vertical transfer [41]. Since our phylogenetic analysis of SN1 was consistent with rRNA data, orthologous SN1 assignment is possible in any vascular plant. Thus, our evolutionary study suggests that MsSN1 from *Medicago sativa* (JQ517286) presents in this work is indeed the ortholog of the StSN1 from *Solananum tuberosum* (Q948Z4), the gibberellin-stimulated transcript 1 (OsGASR1) from *Oryza sativa* (AB192574) and the GAS7 from *Arabidopsis thaliana* (AEC06348) previously described [3,42,43], thus offering a starting point for experimental data integration to in-depth understanding of SN1 function in plants. It is important to note that orthologous identification of snakin-like peptides could not be predicted by comparing their amino acid identity with snakin/GASA-related proteins such as *Arabidopsis* GASAs (Additional file 4).
Therefore, assessment of the correct orthology of snakin/GASA genes requires rigorous phylogenetic analysis.

In order to support the vertical origin of higher plant SN1, the SN1 gene is present in the primitive vascular plant Selaginella moellendorffii (Figure 2). In addition, an exhaustive phylogenetic analysis of proteins within genome databases failed to find orthologous genes of SN1 in the ancestral moss Physcomitrella patens, and in green algae species belonging to the phylum Chorophyta such as Ostreococcus lucimarinus, Micromonas sp. RCC299, Volvox carteri and Chlamydomonas reinhardtii. In agreement with this finding, proteins containing the snakin/GASA domain are widely distributed in land plants but completely absent in moss and green algae (data not shown), suggesting that SN1 is an ancestral gene that appeared in the vascular lineage after the vascular/bryophyte separation. This is not surprising considering that the pioneering ancestors of land plants that dominated terrestrial environments had to confront dramatic stresses, including the infection produced by an extreme diversity of microbial soil communities, selecting a number of genetic innovations [44]. Therefore, the emergence of SN1 could be an adaptation of ancestral plants to land.

Analysis of the in vitro antimicrobial activity of MsSN1

Regarding the in vitro characterization of MsSN1 antimicrobial activity, the MsSN1 gene was expressed in E. coli. The expression of MsSN1 in the transformed E. coli cells was confirmed by RT-PCR analysis (Additional file 5). The antimicrobial activity of MsSN1-free or MsSN1 extracts was tested against the bacterial and fungal alfalfa pathogens Agrobacterium tumefaciens LBA4404 and Phoma medicaginis var. medicaginis CBS 316.90, respectively. Antimicrobial activity of the MsSN1-free extract was undetectable, whereas the MsSN1 extract showed a high inhibitory activity against bacterial growth (Figure 3A-B) and fungal spore germination (Figure 3C-F). As previously described for other members of snakin/GASA peptides [3,4], the antimicrobial activity of MsSN1 against organisms belonging to different domains, at least Bacteria and Eukarya, is congruent with its function as a component of the non-specific immune system of plants.

Analysis of MsSN1 expression

The expression pattern of the MsSN1 gene was analyzed by RT-PCR, to explore the function of MsSN1 in wild type alfalfa plants. MsSN1 expression was detected in all tissues analyzed, including roots, stems, leaves and young floral buds (Additional file 6). RT-qPCR assays were performed to evaluate whether the expression levels of MsSN1 is modified by exposure to microorganisms. MsSN1 transcript levels were about two-fold higher in leaves than in roots when plants were grown under control conditions (Figure 4). Interestingly, MsSN1 levels did not change in response to biotic stimuli, such as the
pathogenic bacterium *A. tumefaciens* LBA4404, the symbiotic strain *Sinorhizobium meliloti* BL225C and the free-living strain *Pseudomonas fluorescens* Pf-5 (Figure 4). In concordance with the characterization of *StSN1* expression in potato [3], the expression pattern of *MsSN1* suggests that *SN1* is a component of the constitutive defense barrier.

**Molecular characterization of transgenic alfalfa plants overexpressing MsSN1**

In order to investigate the antimicrobial activity of *MsSN1* against alfalfa fungal pathogens, transgenic alfalfa plants were generated. The *MsSN1* gene was placed into a constitutive expression cassette under the control of the CaMV 35S promoter (Figure 5A), and introduced in alfalfa plants of the regenerable clone C2-3 by *Agrobacterium tumefaciens*-mediated transformation (Additional file 3).

PCR assays (Additional file 7A) and Southern Blot analyses (Additional file 7B) were used to identify three independent transgenic alfalfa lines, named S1, S2 and S3, harboring the CaMV35S:MsSN1 construct. Evolutionary analyses showed that *Medicago truncatula* and *Lotus japonicus* genomes have only one locus per genome of the SN1 gene, MTR_1g018640 and LJ5G003340, respectively. Consistent with this, as well as with the remarkably conserved genome structure among legumes [45] and the tetrasomic inheritance and outcrossing nature of alfalfa [46], the non-transformed alfalfa plants showed the presence of three copies, probably alleles, of the *MsSN1* gene (Additional file 7B).

As expected, transgenic plants showed extra and differential bands on Southern blot hybridizations, suggesting the incorporation of the transgene in different regions of the alfalfa genome (Additional file 7B). The three events showed stable inheritance of the transgene (data not shown). Expression of the transgene was confirmed by RT-PCR analyses using specific primers designed for the chimeric 5′ UTR (Additional file 7C). Individual lines were also analyzed by RT-qPCR assays for *MsSN1* expression. Transgenic lines showed that the expression level of total *MsSN1* was 100-fold higher than wild type endogenous expression (Figure 5B), suggesting that the three transgenic lines were good candidates to evaluate the effects of overexpressing *MsSN1*.

Similar to transgenic lines from potato, wheat and tomato [6,10,11], overexpression of *MsSN1* did not appear to affect negatively the general phenotype in transgenic alfalfa plants, e.g. there was no reduction in the plant vigor in transgenic plants compared to wild type plants (Additional file 8). Therefore, transgenic alfalfa plants appear to be a suitable platform to study the role and biotechnological potential of *MsSN1*. Unfortunately, no *MsSN1* reduced expression or silenced alfalfa plants were obtained (data not shown). Similar results were
reported by Meiyalaghan [8] where the authors found no regenerant plants using antisense vectors expressing GSL1 (StSN1) and GSL2 (SN2) under their endogenous promoters regulation. These results probably indicate that MsSN1 silencing had detrimental effects on transformation or regeneration efficiency. In concordance with this observation, SN1 silencing drastically affects the cell division and primary metabolism of potato plants [14].

Molecular characterization of two highly virulent alfalfa fungal strains

Anthracnose and spring leaf spot, caused by the fungal pathogens Colletotrichum trifolii and Phoma medicaginis, respectively, are the most destructive infections of alfalfa worldwide. In the analysis of alfalfa resistance to these fungal diseases, there is usually a tradeoff between virulence and characterization of these pathogenic strains. Generally, long-term successive passages of archetype strains in vitro attenuate the virulence of these pathogens in vivo, while pathogens freshly isolated present a high virulence phenotype but their taxonomic position is commonly speculated based on disease symptoms or microscopy observations. In order to bypass this constraint, we used two highly virulent alfalfa fungal strains from alfalfa breeding programs, maintained through sequential plant infections and characterized through amplification, sequencing and phylogenetic analyses of conserved genes and intergenic regions. The evolutionary analyses of ITS region and TUB gene showed that these isolates belong to Colletotrichum and Phoma (Additional file 9). In fact, these DNA fragments from virulent strains (KF846005, KF846006, KF846009 and KF846010) share 100% nucleotide identity with the ITS region and TUB genes from C. trifolii CBS 158.83 (KF178478 and KF178599) and P. medicaginis CBS 316.90 (GU237828 and GU237630), respectively. In accordance with these data, these isolates were named Colletotrichum trifolii strain CT2 and Phoma medicaginis strain CT1. Unlike the well-characterized fungal strains C. trifolii CBS 158.83 and P. medicaginis CBS 316.90, strains C. trifolii CT2 and P. medicaginis CT1 are highly virulent on alfalfa and infect a wide range of commercial cultivars as well as the alfalfa clones used in this work, C2-3 and 432-19-17. This confirms that the form of conservation of fungal pathogens is essential to maintain their alfalfa virulence, and therefore, to study the genetic basis of resistance to fungi in alfalfa.

Antifungal in vitro activity of MsSN1-overexpressing transgenic plants

As a first step to investigate the antifungal activity of transgenic alfalfa lines, in vitro challenge assays were performed. Detached leaflets were inoculated with a spore...
suspension of two virulent and molecularly characterized alfalfa fungal strains *P. medicaginis* CT1 and *C. trifolii* CT2 incubating the leaves in petri plates for 7 days (Figure 6A-B). Disease severity was estimated by visually scoring disease symptoms in infected leaflets as a percentage of leaflets showing disease lesions (Figure 6C). The three transgenic lines (S1- S3) showed significant lower percentage of diseased leaflets than wild type plants when challenged with both *P. medicaginis* CT1 (Figure 6A) and *C. trifolii* CT2 virulent strains (Figure 6B), suggesting an antimicrobial function of MsSN1 under biotic stress conditions. Moreover, this small-scale study could facilitate rapid visual screening of a large number of transgenic events (Figure 6D), saving time, reducing cost, and speeding up the introgression of this antimicrobial transgene into commercial varieties, especially in outcrossing species such as alfalfa [47].

**Antifungal in vivo activity of MsSN1-overexpressing transgenic plants**

The three transgenic alfalfa lines (S1-S3) were chosen for progeny analysis of their potentially enhanced biotic resistance. The clone 432-19-17 was used as a pollen donor in a sexual cross with the transgenic lines to avoid the common inbreeding depression of progeny from self-crossed alfalfa plants. The presence and expression of the transgene in leaves of the progeny plants were assayed by PCR and RT-qPCR analyses, respectively (data not shown). **In vivo** assays for antifungal activity against *P. medicaginis* CT1 were carried out spraying a spore solution to transgenic and wild type plants. Plant disease severity was estimated by evaluating agronomic parameters for two months. Wild type plants manifested all the symptoms of the disease [48,49], including dark brown lesions on the stems, dark brown lesions on the leaflets that begin with specific dark color injuries, wilting leaves, chlorosis, and defoliation. Transgenic lines presented scarce symptoms and usually looked like control plants (data not shown).

Firstly, leaflet disease was estimated by visually scoring disease symptoms in infected plants. One month post-infection, the percentage of diseased leaflets was significantly lower in transgenic lines than in wild type plants (Figure 7A). Two months post-inoculation, disease severity was analyzed by evaluating the vigor of the plants, the number of plants with regrowth and the percentage of highly defoliated plants. Transgenic plants showed significantly higher levels of regrowth than wild type plants (Figure 7B, F and G). In fact, disease symptoms were present in wild type regrowth but not in transgenic plants (Figure 7G). Overexpression of MsSN1 in alfalfa improves plant vigor under fungal stress conditions (Figure 7C). In addition, wild type plants showed more defoliation than transgenic plants but the differences

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**Figure 6** Antifungal in vitro activity of MsSN1 overexpressing transgenic plants. (A) Diseased leaflets related to leaflets inoculated with *Phoma* (#diseased leaflet/#total leaflet). (B) Diseased leaflets related to leaflets inoculated with *Colletotrichum* (#diseased leaflet/#total leaflet). (C) Damage score: 1. Healthy leaflet, 2. Countable injuries, 3. Uncountable injuries, 4. Chlorosis, 5. completely damaged. (D) Representative *Phoma* assay. Wt: wild type. S1-S3: MsSN1 transgenic plants. All values are log means ± SEM (n = 10–30). Asterisks indicate a statistically significant difference (Turkey: "p < 0.5; **p < 0.01 ***p < 0.001). Leaflets were considered diseased when they showed symptoms 3, 4 or 5 in the damage score. Disease severity was estimated from scoring 30 to 60 detached leaflets from three individual plants.
were not statistically significant (Figure 7D). Altogether, transgenic plants showed a better performance against Phoma infection than wild type plants, suggesting an antifungal activity of MsSN1 in vivo.

**Coevolution evidences of MsSN1 and rhizobia**

Similar to observations on other effector molecules of innate immunity [50], constitutive expression of MsSN1 could provide alfalfa a selective advantage over the pathogens during the early stages of microbial infection. However, it can also have adverse effects on beneficial microorganisms such as nitrogen-fixing symbiotic bacteria. To investigate this apparent paradox, the *in vitro* antimicrobial activity of MsSN1 against nine rhizobacterial strains isolated from different legume plants and soil were analyzed. All rhizobacteria isolated from alfalfa nodules (*Sinorhizobium meliloti* BL225C, *Sinorhizobium meliloti* SM11 and *Rhizobium* sp. Or 191) showed no inhibition against the MsSN1 extract (Table 1). In contrast, the symbiotic nitrogen fixation strains isolated from other legume species, such as medick (*Sinorhizobium medicae* WSM419), soybean (*Sinorhizobium fredii* USDA 257 and *Bradyrhizobium japonicum* USDA110), bean (*Rhizobium etli* CFN 42) and Lotus spp. (*Mesorhizobium loti* MAFF303099),...
were susceptible to the MsSN1 extract (Table 1). These results suggest that alfalfa and their symbiotic bacteria may have coevolved.

There is a fine line between bacterial symbiosis and chronic infection. While one is beneficial the other is detrimental. Recent findings suggest that both share mechanisms for sidestepping host defenses [50]. Legumes evolved rapidly and shortly after their origin, and nodulation most likely evolved several times during their divergence [51]. These plants have symbiotic nitrogen-fixing bacteria living in root nodule compartments that also contain antimicrobial compounds. To avoid infection with phytopathogenic bacteria, nitrogen-fixing rhizobial bacteria and leguminous plants have developed complex signal exchange mechanisms that allow a specific bacterial strain to induce its specific host plant species to form invasion structures through which the bacteria can enter the plant root [52]. The evidence of co-evolution presented here suggested that MsSN1 peptide could be part of the battery of compounds involved in discriminating the symbionts from other microbes, so as to allow the beneficial infection and inhibit colonization of potentially pathogenic microbes.

In concordance with the hypothesis of plant-microbe coevolution, MsSN1-overexpressing transgenic plants showed nodule production of *S. meliloti* BL225C comparable with that of the wild type plants (Figure 8A-B), suggesting that MsSN1 does not affect plant infection when nitrogen-fixing bacterial strains have an alfalfa origin. Then, to explore the effect of MsSN1 on free living strains of rhizobacteria as *Pseudomonas*, the antimicrobial activity of MsSN1 on *P. fluorescens* Pf-5 isolated from soil samples was studied. Interestingly, *P. fluorescens* Pf-5 showed hypersensitivity to the MsSN1 extract.

| Strain                        | Family           | Source of the strain | Zone of inhibition (mm) |
|-------------------------------|------------------|----------------------|-------------------------|
| *Bradyrhizobium japonicum* USDA110 | Bradyrhizobiaceae | Soybean              | 1.38 ± 0.13             |
| *Mesorhizobium loti* MAFF303099 | Phyllobacteriaceae | Lotus spp.            | 1.50 ± 0.30             |
| Rhizobium sp. Or 191          | Rhizobiaceae     | Alfalfa              | 0                       |
| Rhizobium etli CFN 42         | Rhizobiaceae     | Bean                 | 1.18 ± 0.09             |
| *Sinorhizobium meliloti* BL225C | Rhizobiaceae     | Alfalfa              | 0                       |
| *Sinorhizobium meliloti* SM11 | Rhizobiaceae     | Alfalfa              | 0                       |
| *Sinorhizobium medicae* WSM419 | Rhizobiaceae     | Medick               | 1.26 ± 0.06             |
| *Sinorhizobium fredii* USDA 257 | Rhizobiaceae    | Soybean              | 1.13 ± 0.13             |

Figure 8 Bacterial colonization of MsSN1-overexpressing transgenic plants. Analysis of alfalfa-*Sinorhizobium meliloti* BL225C interactions: number of nodules in two-month-old alfalfa plants. (A) Photo of nodulated root. (B) Study of alfalfa colonization by *Pseudomonas fluorescens* Pf-5. (C) MsSN1-overexpressing plants. Wt: wild type. All values are means ± SEM (n = 20). N.S: non-significant, ***p < 0.001, t-test.
(Zone of inhibition = 1.76 ± 0.06 mm). In agreement with the in vitro activity of MsSN1, P. fluorescens Pf-5 showed differential proliferation in the rhizosphere of wild type and transgenic lines of alfalfa 30 days after inoculation, suggesting an increased antibacterial activity in MsSN1-overexpressing plants (Figure 8C). These results show additional in vitro and in vivo evidence of MsSN1 as a part of the innate immune response and the wide range of activity of MsSN1.

Conclusions
Previous studies and the results reported here show that snakin peptides have important and ancestral roles in land plant innate immunity. Interestingly, MsSN1-overexpressing alfalfa transgenic plants show increased antimicrobial activity against virulent fungal strains without altering the nitrogen-fixing symbiosis, opening the way to the production of effective alfalfa transgenic cultivars for biotic stress resistance. In this work, data also suggest a coevolutionary process, in which alfalfa exerts a selection pressure for resistance to MsSN1 on rhizobial bacteria.

Additional files

Additional file 1: Table of primers used in this work.
Additional file 2: Similarity of the product of MsSN1 gene to other snakin/GASA proteins from plants.
Additional file 3: Alfalfa transformation.
Additional file 4: Identity of the product of MsSN1 gene to other snakin/GASA proteins from Arabidopsis.
Additional file 5: RT-PCR studies of MsSN1 gene expression in E. coli pS333 (1) and E. coli pS333-MsSN (2) strains.
Additional file 6: RT-PCR assays showing MsSN1 expression in young floral buds (A-B), leaves (C-D), stems (E-F) and roots (G-H).
Additional file 7: Molecular characterization of transgenic alfalfa lines overexpressing MsSN1.
Additional file 8: Vigor phenotype of transgenic (S1) and wild type (Wt) alfalfa plants.
Additional file 9: Phylogenetic analysis of fungal strains isolated from alfalfa.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
Conceived and designed the experiments: GCS; NDA; JPM; MEP. Performed the experiments: ANG; NDA; GCS; ARF; MCG; MJD. Analyzed the data: GS; NDA. Wrote the paper: GS; NDA; CAB. All authors read and approved the final manuscript.

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