Metabolomic analysis of cold acclimation of Arctic Mesorhizobium sp. strain \(N_{33}\)
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Introduction

Bacteria have developed many strategies at the transcriptional and post-transcriptional levels to enhance their abilities to withstand cold temperature stress. Some molecular responses to stress are general while others can be specific [1]. For instance, by using genome, cell physiology and transcriptome analyses, it has been shown that at −15°C, the permafrost bacterium Planococcus halocryophilus strain Or1, specifically regulates genes encoding for: the conversion of saturated fatty acids to unsaturated and branched fatty acids; the remodeling of cytoplasmic membrane and cell envelope features; transport systems, chaperone proteins, accumulation of cryoprotectant compounds, and transcriptional regulation [2]. The specific cold adaptive features in bacteria include global resource efficiency, amino acid substitution in cold-loving bacteria, the synthesis of lipid desaturases has also been detected in strain TAC125. These enzymes increase membrane fluidity, and protect the cell against dioxygen and detoxify the cells at low temperature. On the other hand, as a general response, housekeeping genes are also regulated by a variety of stressors including low temperature which allows these bacteria to down-regulate their metabolism to optimize general cell functions in order to withstand cold stresses. Other general response mechanisms include higher turnover of macromolecules, tighter maintenance of intercellular pH, greater osmotic regulation, motility, stopping biomass production and decreasing the activation energy before a pre-exponential growth phase [6]. Reduction of growth and suppression of the genes involved in translation and ribosomal biogenesis in E. coli have been observed and can be considered as a general response and an important strategy under stress conditions [7]. Transcriptional analysis of the Arctic Mesorhizobium N33 revealed the down-regulation of many housekeeping genes that encode for the cell envelope and outer membrane biogenesis.
functions, as well as cell motility and secretion at low temperature (unpublished data).

Studies on cold adaptations of the mesophilic [8–10] and psychrophilic bacteria [11,12] indicated that they are widely heterogeneous in their genomes content and encompass broad ranges of complex network strategies to survive at low temperature. Most cold adaptation comprehensive studies were mainly performed by applying high-throughput genome sequencing [4,5,11] and other omics technologies such as proteomics [13,14] and transcriptomics [15]. In E. coli, it has been shown that under different perturbations (cold, heat, lactose diauxie, and oxidative stress) the metabolite profiles are much more stress-specific when compared with transcriptomic changes [16].

In our studies, the transcriptional analysis of Arctic Mesorhizobium N33 at different cold temperature treatments has been informative (unpublished data). For instance, we observed that some amino acids, polyunsaturated fatty acids, and cryoprotectants changed significantly. Since transcriptomics studies only allow the evaluation of molecular adaptation mechanisms at the level of gene expression and did not reflect much higher specificity during early stress adaptation [16], we hypothesized that using metabolomics measurements will allow a better understanding of N33 cells response to its environmental changes [16–18]. Furthermore, metabolites are the end products of cellular processes, and they can be considered as a link between genotype (gene function) and phenotype [19]. It was previously shown that metabolic profiles could be good and specific indicators for monitoring cellular responses to biotic and abiotic stresses [18]. Nevertheless, our understanding of the metabolite content of cold adapted bacteria is very limited.

The psychrotrophic Arctic Mesorhizobium sp. N33 is the best known cold-adapted symbiotic N2-fixing rhizobium isolated from nodules of Oxytropis arctica, an indigenous legume from Canada’s eastern Arctic [20]. Strain N33 has an intrinsic resistance to streptomycin and it has been conformable to genetic manipulations [21–23]. Strain N33, like its closely related strain N31, has a growth temperature range between 0°C and 30°C, and can establish an efficient symbiosis with the temperate forage crop enzyme (catalyzing the reduction of N2) and other omics technologies such as proteomics [13,14] and transcriptomics [15]. In E. coli, it has been shown that under different perturbations (cold, heat, lactose diauxie, and oxidative stress) the metabolite profiles are much more stress-specific when compared with transcriptomic changes [16].

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adaptations used by the Arctic Mesorhizobium N33 at low temperature.

Results and Discussion

Experimental Design and Metabolic Profiles

GC-MS and NMR were used for the identification and quantification of water soluble metabolites and fatty acids in cells of Mesorhizobium N33 growing at or exposed to, suboptimal temperatures as compared to cells grown at 21°C considered as control in this study. Water-soluble metabolites and lipids were extracted by phase separation using a biphasic system as described in the materials and methods section. NMR analyses of water soluble metabolites allowed the identification and quantification of 29 compounds (Table 1, and Table S1). With GC-MS, 26 water-soluble metabolites were identified, of which nine were also detected by NMR (Table 1 and Figure 1).

Mean concentrations (µM) of water-soluble metabolites (measured by NMR) and their standard deviation (SD) are shown in the supporting information (Table S1). Fatty acids from total, neutral, glyco- and phospho- lipids were identified and quantified using GC-MS. We were able to identify and quantify 13 types of fatty acids from total lipids, 17 from neutral lipids, 17 from phospholipids, and 17 from glycolipids. The complete list of fatty acids, mean concentrations (expressed as mole % of total fatty acids) and their standard deviation (SD) are shown in supporting information (Tables S2, S3, S4, and S5). A list of the 20 fatty acids identified in N33 is presented in Table 2.

Multivariate Statistical Analyses of Water Soluble Metabolites

Principal components analysis-PCA (Figure 2A) and heatmap visualization (Figure S2) show that growth temperature significantly affects water soluble metabolites in Mesorhizobium N33. Cells grown at 10°C had water soluble metabolites contents, comparable to that of the control cells grown at 21°C. Nevertheless, slight differences in the level of metabolites accumulation were observed between the two temperatures. Cells grown at 4°C clustered in a distinct group indicating that at this suboptimal temperature, changes in metabolites levels (up and down) were more pronounced. Out of 29 water soluble metabolites, 19 showed a high accumulation in at least two independent biological repetitions and 10 compounds were suppressed during acclimation to 4°C. Isobutyrate, sarcosine, threonine, and valine were up-regulated in all three biological replicates at 4°C (Figure S2), suggesting that these compounds might be required for energy conservation [2,16]. The significant compounds of N33 water soluble metabolites after different times of exposure to suboptimal

![Figure 1. Venn diagram showing the overlap of Mesorhizobium N33 water soluble metabolites detected by NMR and GC-MS, and fatty acids from different lipids detected by GC-MS.](https://www.plosone.org/article/static/doi:10.1371/journal.pone.0084801.g001)
4°C temperature (T1 to T5; 2 to 240 min), are shown based on PLS-DA (partial least squares discriminate analysis) (Figure 2B) and heatmap visualization (Figure S3). The significant (P<0.01) permutation test confirmed the PLS-DA analysis. The optimal PLS-DA model for water soluble metabolites data (measured by NMR) used the top four components with a $Q^2 = 0.78$. Results revealed that some metabolites accumulated while others decreased after different times of exposure to 4°C compared to the control (T0 = 21°C). The most important compounds are indicated in Figure S2 and they include: sarcosine, glycine, lactate, glutamate, glucose, methionine, 3-hydroxybutyrate, acetate, tyrosine, and isoleucine. The increase in amino-acid levels might result, at least in part, from an increase in protein degradation [32] necessary for the elimination of abnormal proteins resulting from stress conditions. Abundance of the amino-acids might also be due to the demand for the synthesis of new important proteins (chaperones) under stress conditions [16,33]. A genomic study has supported the thermal flexibility apparent from amino acid distribution in the permafrost bacterium (Planococcus halocryophilus strain Or1) [2].

Multivariate Statistical Analyses of Fatty Acids from Neutral Lipids

PCA analysis (Figure 3A) and heatmap visualization (Figure S4) indicated that neutral lipids in cells of Arctic strain N33 grown at 10°C exhibited fatty acids content comparable to the one observed when cells were grown at 21°C. However a distinct content was obtained when N33 was cultivated at the suboptimal 4°C temperature. Out of 17 measured fatty acids, linoleic acid 18:2(6,9) showed higher accumulation in cells grown at 4°C. Many metabolites have shown same change trends at 10 and 21°C (Figure S4). The significant fatty acids compounds from neutral lipids after different times of exposure to suboptimal 4°C temperature, are shown according to PLS-DA analysis (Figure 3B), and heatmap visualization (Figure S5). The permutation test was statistically significant (P<0.07) which confirmed the results of PLS-DA analysis. The optimal PLS-DA model for fatty acid of neutral lipids used the top two components with a $Q^2 = 0.37$.

### Table 1. Water soluble metabolites of Arctic *Mesorhizobium* N33 detected by NMR and by GC-MS.

| Metabolites identified by NMR | HMDB ID | Metabolites identified by GC-MS | HMDB ID |
|------------------------------|---------|--------------------------------|---------|
| 3-Hydroxybutyrate            | HMDB00357 | 2,3-Dimethoxy phenylpyruvic acid | HMDB00205 |
| Acetate                      | HMDB00042 | 4-aminobutyric acid            | HMDB00112 |
| Acetone                      | HMDB01659 | Beta-Amino isobutyric acid     | HMDB02166 |
| Alanine                      | HMDB00161 | Beta-Hydroxybutyric acid       | HMDB01873 |
| Choline                      | HMDB00097 | Adenosine                      | HMDB00050 |
| Ethanol                      | HMDB00108 | Alanine                        | HMDB00161 |
| Formate                      | HMDB00142 | Arabin-Hexos-2-ulos            | CHEBI:27973 |
| Glucose                      | HMDB00122 | Arabinofuranose                | HMDB12325 |
| Glutamate                    | HMDB03339 | Cadaverine                     | HMDB02322 |
| Glycerate                    | HMDB00131 | Galactose                      | HMDB00143 |
| Glycine                      | HMDB00123 | Gluconic acid                  | HMDB00625 |
| Isobutyrate                  | HMDB01873 | Glucose                        | HMDB00122 |
| Isoleucine                   | HMDB00172 | Glycerol                       | HMDB00131 |
| Lactate                      | HMDB00190 | Lysine                         | HMDB00182 |
| Leucine                      | HMDB00687 | Malonic acid                   | HMDB00691 |
| Lysine                       | HMDB00182 | Mannitol                       | HMDB000765 |
| Malonic acid                 | HMDB00691 | Mannose                        | HMDB00169 |
| Mannitol                     | HMDB00765 | Methionine                     | HMDB00696 |
| Methionine                   | HMDB00696 | Methylaminobutyric acid        | HMDB00039 |
| N-Acetylglycerine            | HMDB00532 | Methylmalonate                 | HMDB00202 |
| N-Carbamoyl-β-alanine        | HMDB00026 | Oxalic acid                    | HMDB02329 |
| Oxypurinol                   | HMDB00786 | Phenylalanine                  | HMDB00159 |
| Phenylacetate                | HMDB00209 | Putrescine                     | HMDB01414 |
| Sarcosine                    | HMDB00271 | Succinic acid                  | HMDB00254 |
| Succinate                    | HMDB00254 | Tyrosine                       | HMDB00158 |
| Threonate                    | HMDB00943 | Valine                         | HMDB00883 |
| Threonine                    | HMDB00167 |                                |          |
| Tyrosine                     | HMDB00158 |                                |          |
| Valine                       | HMDB00883 |                                |          |

Compounds indicated in bold were detected by NMR and GC-MS. HMDB: Human metabolome database (http://www.hmdb.ca/).

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Table 2. Fatty acids identified by GC-MS in cold exposed Arctic Mesorhizobium N33.

| Fatty Acids | Common Name          | IUPAC Name (Systematic)          |
|------------|----------------------|----------------------------------|
| C12        | Lauric (Dodecanoic)  | Dodecanoic acid                  |
| C14        | Myristic (Tetradecanoic) | Tetradecanoic acid               |
| C14:1(11)  | Myristoleic acid     | Tetradecenoic acid               |
| C15        | Pentadecanoic acid   | Pentadecanoic acid               |
| C16        | Palmitic acid        | Hexadecanoic acid                |
| C16:1(7)   | Palmitoleic acid     | Hexadecenoic acid                |
| C16:1(9)   | Palmitoleic acid     | Hexadecenoic acid                |
| C18        | Stearic acid         | Octadecanoic acid                |
| C18:1      | Oleic acid           | Octadecenoic acid                |
| C18:1(10)  | Oleic acid           | Octadecanoic acid                |
| C18:1(9)   | Oleic acid           | Octadecanoic acid                |
| C18:2(6,9) | Linoleic acid        | Octadecadienoic acid             |
| C18:2(9,12)| Linoleic acid        | Octadecadienoic acid             |
| C19        | Nonadecanoic acid    | Nonadecanoic acid                |
| C19:1(10)  | Nonadec-10(Z)-enoic | Nonadecenoic                     |
| C20        | Arachidic acid       | Eicosanoic acid                  |
| C20:1      | 11-Eicosenoic acid   | Eicosenoic acid                  |
| C20:1(11)  | 11-Eicosenoic acid   | Docosanoic acid                  |
| C22:1      | Erucic acid          | Eicosanoic acid                  |
| C22:1(13)  | Cetoleic acid        | Docosanoic acid                  |

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Multivariate Statistical Analysis of Phospholipids Fatty Acids Content

The PCA analysis (Figure 4A) of the phospholipids fatty acids content of N33 growing at constant temperatures and heatmap visualisation (Figure S6) show distinct trend of metabolite changes at each growth temperature tested (21°C, 10°C and 4°C). Out of 17 measured fatty acids from phospholipids, the linoleic acids 18:2(9,12), 18:2(6,9), palmitoleic acid 16:1(7), and cetoleic acid 22:1(13) showed higher accumulation in N33 growing at 4°C (Figure S6). Significant changes in phospholipids fatty acids after different times of exposure to suboptimal 4°C temperature revealed by PLS-DA analysis (Figure 4B), and heatmap visualisation are shown in Figure S7. The significant permutation test (P<0.01) confirms the results of PLS-DA analysis. The optimal PLS-DA model for phospholipids fatty acids used the top five component with a Q² = 0.74. The PLS-DA and heatmap show increases or decreases in metabolite after each time of exposure (from T0 to T5) to 4°C.

Multivariate Statistical Analysis of Fatty Acids from Glycolipids

The PCA analysis (Figure 5A) of the glycolipids fatty acids present in N33 growing at constant temperatures and heatmap visualisation (Figure S8) show different trends of metabolic changes. N33 cells growing at 4°C show a distinct trend as compared to cells cultivated at 21°C, whereas an intermediate level of metabolite changes is observed with cells growing at 10°C. Out of 17 measured fatty acids from glycolipids, linoleic acid 18:2(6,9), nonadecanoic acid (C19), myristic (tetradecanoic) acid (C14) concentration increased at 4°C, and linoleic acid 18:2(6,9) increased at 10°C. These fatty acids were present at lower concentrations in cells cultivated at 21°C (Figure S8). The significant fatty acids compounds of glycolipids after different times of exposure to suboptimal 4°C temperature identified by PLS-DA analysis (Figure 5B) and heatmap visualization (Figure S9). The significant permutation test (P<0.01) confirms the results of PLS-DA analysis. The optimal PLS-DA model for fatty acid from glycolipids used the top three component with a Q² = 0.75. The PLS-DA and heatmap showed that time of exposure to suboptimal 4°C significantly influenced the concentration of glycolipids fatty acids in the arctic strain N33 (Figure 5B).

Univariate Statistical Analyses of Fatty Acids from Total Lipids

The PCA analysis (Figure 6A) and heatmap visualisation (Figure S10) show that the total fatty acids content of N33 is significantly affected by growth temperature. Distinct trends of metabolic changes at 21°C, 10°C and 4°C were observed by heatmap visualization (Figure S10). Out of 13 total fatty acids, at least 6 compounds showed accumulation at 4°C and 7 others decreased. These patterns were opposite in N33 growing at 21 and 10°C. Significant fatty acids from total lipids after different times of exposure to suboptimal 4°C temperature revealed by PLS-DA analysis (Figure 6B), and heatmap visualisation is shown in Figure S11. The significant permutation test (P<0.01) confirms the results of the PLS-DA analysis. The optimal PLS-DA model for fatty acids from total lipids used only the top component with a Q² of 0.31. The PLS-DA and heatmap showed different trends of metabolic changes for each time of exposure to 4°C.

Unsupervised PCA analysis of fatty acids from the 4 lipid classes, disclosed that the metabolite profiles of N33 cells grown at 21°C, 4°C or 10°C are each clustered into a distinct group, contrary to cells grown at 21°C and exposed to 4°C for 2 to 240 minutes, thereby suggesting a specific metabolic acclimation to cold.

To distinguish and better understand the detailed information arising from the multivariate analyses, and to identify metabolite features that are significantly different between each treatment and the control, univariate statistical analysis (i.e. t-tests, determining the fold change (FC) and volcano plots) were performed. These analyses were applied to five different metabolite data sets (total fatty acids, and fatty acids from neutral, glyco- and phospholipids, and water soluble compounds) and are presented below.

Univariate Statistical Analyses of Fatty Acids

When grown at 21°C the arctic Mesorhizobium N33 had a total fatty acid content formed mainly by the following lipids (mole % of total lipids): 18:1 (54%); 16:0 (26%); 18:0 (13.5%) and 19:1(10) (3.19%) (Table S2). Monounsaturated 18:1, along with the saturated 16:0 and 18:0 fatty acids are the major lipids found in rhizobia grown at optimal temperature [34–37]. The fatty acids identified by GC-MS which showed significant (P<0.05; FC ≥2) concentration shifts at low temperatures are shown in Figure 7.

The maximum increases in fold-change (FC) were observed when N33 was grown at 4°C or 10°C (GT4 or GT10). In fact, when grown at 4°C and 10°C, N33 contained respectively 18 and 17 times more linoleic acid 18:2 (6, 9) than cells grown at 21°C. The concentration of the fatty acid 19:1(10) significantly decreased in cells grown at 4°C. N33 cells growing at 21°C (T0) then exposed to 4°C for 2 min (T1) did not exhibit any significant change in fatty acids measured from total lipids. However, when exposed to cold for 4 or 8 min (T2 and T3), 5.6 and 2.2 fold increases were observed for fatty acids 22:1 and 19:1(10) (Figure 7A).

Univariate statistical analysis indicated that the concentration of 13 [5 saturated and 8 unsaturated] fatty acids from neutral lipids
A: PCA analysis was performed based on 29 water soluble metabolites. B: PLS-DA plot showing grouping of compounds (permutation test, P < 0.01) according to the different time of exposure to a suboptimal 4°C temperature. T0 = 21°C (reference), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. The optimal PLS-DA model for water soluble metabolites data (measured by NMR) used the top four components with a $Q^2 = 0.78$. doi:10.1371/journal.pone.0084801.g002

significantly (P ≤ 0.05, FC ≥ 2) changed in N$_{33}$ cells submitted to cold treatments (Figure 7B). Overall, 12 accumulations and 13 reductions in fatty acids from neutral lipids were observed with all cold treatments. As observed with fatty acids of total lipids, at 4°C N$_{33}$ cells contained more (FC = 35) fatty acid 18:2 (6, 9) than cells of the control (growth at 21°C). Comparable results were observed at 10°C (FC = 6). An important increase (FC = 45) in the concentration of fatty acid 14:1 (11) was observed only in N$_{33}$ cells exposed to 4°C for 1 h. Under the same condition, the highest fold change ratio (FC = 9) was observed for the saturated fatty acid 19:0. The most important decrease in concentration was observed for the saturated fatty acid 20:0 (FC = 0.27), in N$_{33}$ cells after 8 min exposure to 4°C (Figure 7B).

The significant changes in fatty acids from glycolipids of Mesorhizobium N$_{33}$ exposed to cold temperature are shown in Figure 7C. Out of the 17 fatty acids detected from glycolipids (Figure 1 and Table S4), 4 saturated and 6 unsaturated changed concentrations after cold treatments. Significant (P ≤ 0.05, FC ≥ 2) levels of accumulation resulting from exposure to cold, ranged from 2.32 FC for fatty acid chain 22:1 (13) at 4°C to 12.5 FC for fatty acid 18:2 (6, 9) in N$_{33}$ growing at 10°C.

Reduction in the concentration of fatty acids from glycolipids in N$_{33}$ ranged from a FC of − 38.46 for fatty acid 19:1 (10) at 4°C to a FC of −2.22 for fatty acid 18:0 for growth at 10°C. As revealed by univariate statistical analysis 8 (5 unsaturated and 3 saturated) out of 17 fatty acids from phospholipids identified in Mesorhizobium strain N$_{33}$ (Figure 1 and Table S5), showed significant (P ≤ 0.05, FC ≥ 2) changes in concentrations after cold perturbations. N$_{33}$ cells grown at 4°C or at 10°C, contained respectively 24.5 and 16.6 times more linoleic acid 18:2(6, 9) than cells growing at 21°C (Figure 7D). Phospholipids from strain N$_{33}$ cultivated at 4°C, contained substantially more fatty acids 16:1(7) (FC = 3.8), 14:0 (FC = 3.9), and 18:2(9, 12) (FC = 11.6) than the control. In N$_{33}$ cells exposed to 60 min 4°C an important increase in fatty acids 14:0 (FC = 10.4) and 14:1 (11) (FC = 44.7) were observed (Figure 7D). The concentration of fatty acid 19:1(10) in N$_{33}$ significantly decreased in cells cultivated at 4°C (FC = −40) and at 10°C (FC = −5.55; Figure 7D).

This study shows that some saturated and many unsaturated fatty acids of the Arctic Mesorhizobium N$_{33}$ were affected by suboptimal temperatures. Saturated fatty acids have the least steric interference with neighboring methylene groups, and so they may be used to enhance the rigid structure of the membrane and protect the cell conformation. Protection of psychrophilic bacteria from cold temperatures by the overproduction of polyunsaturated fatty acid was previously demonstrated by using transcriptional and biochemical analyses [38,39]. Unsaturated fatty acids have kinks which limit acyl-chain packing and cause a melting point reduction (melting point for linoleic acid 18:2(9, 12) is −8.5°C). This feature improves cell membrane fluidity at low temperatures and the increased production of unsaturated fatty acids appears to be one mechanism for the low temperature adaptation in Arctic Mesorhizobium N$_{33}$. Our results corroborate previous observations indicating that rhizobia increase the production of unsaturated fatty acids during growth at cold temperatures [35,37].

Interestingly, a marked effect of temperature on the fatty acid composition in N$_{33}$ was reflected by the different trends observed, when cells were grown at 4°C or 10°C (Figures 3A to 6A).
Nevertheless, at both temperatures, N33 tends to produce significantly more linoleic acid 18:2 (6, 9) and less of 19:1(10) than the control cells for the different types of fatty acids studied. Changes in fatty acid composition following different time of exposure to cold temperatures were more diverse.

Univariate Statistical Analysis of Water Soluble Metabolites

The significant changes (P ≤ 0.05, ratio ≥ 2) in the water-soluble metabolites content of Mesorhizobium N33 subjected to different suboptimal temperature treatments were identified by NMR and are summarized in Figure 8. Univariate statistical analysis revealed that the concentration of several compounds were significantly (P ≤ 0.05, ratio ≥ 2) altered by cold. In comparison to cells grown at 21°C, N33 grown at 4°C contained significantly more valine, threonine, sarcosine and isobutyrate with FC ranging from 2 to 19.4 (Figure 8 and Table S1). Seven water soluble compounds showed a significant concentration decrease at 4°C, but the most important were observed with 3-hydroxybutyrate and oxypurinol. Fewer changes were observed in N33 grown at 10°C compared to 21°C. In fact only 3 compounds changed concentrations and N-carbamoyl-β-alanine displaying the lowest FC (Figure 8).

The metabolite with the highest fold change for all temperature conditions was isobutyrate, with a 19.4 fold change increase found in N33 growing at 4°C (GT4). The greatest reduction in water-soluble metabolites was observed at some conditions with a 0.003 fold change. This lowest level of metabolites fold change might be either the result of being below the instrument detection limit or that the level of the metabolite production has been very low compared to control conditions (N33 grown at 21°C). As a general rule, NMR methods are not particularly sensitive, but using this technique reduces the loss of compounds that may occur during sample preparation [29]. Combining GC-MS and NMR can compensate for the lack of coverage of each platform, but both are still insufficiently sensitive to cover all metabolites. It has been suggested that LC-MS or DI-MS (direct injection of metabolites spectrometry) might be the best methods for metabolomics because of their high sensitivity despite their bias against hydrophilic metabolites [26].

Overall, our metabolomic study of the Arctic Mesorhizobium N33 using GC-MS and NMR was able to identify 110 compounds involved in central carbon metabolism, essential biosynthetic pathways, secondary metabolism and lipids under different low temperature treatments. GC-MS could measure 64 fatty acids (Figure 1, Tables S2, S3, S4 and S5) and identify a variety of amino acids and organic acids (Table 1). NMR spectroscopy provided complementary information by enabling the quantification of 29 water-soluble metabolites (Figure 1, Table 1).

The metabolic analysis of Arctic strain N33 indicated that among the lipid-soluble compounds, poly-unsaturated linoleic acids 18:2(9, 12) and 18:2 (6, 9) were the most abundant fatty acids present in cells grown at 4°C or 10°C, as compared to the control cells growing at 21°C. The mono-unsaturated fatty acid (myristic acid) 14:1(11) from phospho- and neutral lipids was the most

Figure 3. PCA and PLS-DA of fatty acids from neutral lipids present in Mesorhizobium N33 growing at constant temperatures or exposed to suboptimal 4°C. Growth temperatures: GT21 = 21°C (control); GT4 = 4°C; GT10 = 10°C. For all data, row-wise normalization was realized with a pooled averaged reference sample (GT21 or T0), and data were auto scaled and log transformed. A: PCA analysis was performed based on 17 fatty acids. B: PLS-DA plot showing compound changes (permutation test significant at P < 0.01) after exposure for different time to a suboptimal 4°C temperature. T0 = 21°C (reference), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. The optimal PLS-DA model for fatty acids from neutral lipids used the top two component with a Q² = 0.37.

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significantly overexpressed (45-fold change) after exposure for 60 min to 4°C. These fatty acids are known to provide physical membrane flexibility adaptation and to supply energy to cells [40]. Analysis of water-soluble compounds revealed that isobutyrate, sarcosine, threonine and valine increased during growth at 4°C and after exposure for different times to 4°C of cells initially grown at 21°C (T0). Among the water-soluble metabolites, isobutyrate was highly upregulated (19.4-fold) in cells grown at 4°C, suggesting that this compound is a precursor for the cold-regulated fatty acid modification to low temperature adaptation [41]. We have observed that some metabolites decreased in N33 under cold conditions. This might be caused by growth cessation or reduction, which is an important strategy to adjust cellular physiology to cold stresses [16].

Sarcosine (N-methylglycine) is an intermediate and a by-product of glycine synthesis and degradation. Sarcosine oxidase demethylates sarcosine to glycine [42]. It can be derived from the catabolism of betaine, and metabolism of choline, betaine and sarcosine may be linked together [4]. In addition, sarcosine has a significant role in the glutathione metabolic pathway and is a cryoprotectant in psychrophilic bacteria grown at low temperature [5]. These metabolites might have a substantial impact in conferring cold resistance ability to the strain N33 at 4°C by potentially acting as cryoprotectants. Accumulation of cryoprotectants and the production of unsaturated and short chain fatty acids are considered examples of specific metabolic responses to low temperatures in psychrophilic bacteria as previously shown by transcriptomic and genomic sequencing [4]. Sarcosine is also involved in low temperature osmoadaptation [43] and can be used as a source of carbon, nitrogen and energy [4]. Genome sequencing analysis of the psychrophilic bacterium Colwellia psychrerythraea 34H using genomic and proteomic methodologies [4], and phylogenetic diversity and metabolic potential in a glacier metagenome study revealed that many genes are involved in the synthesis of cryo-osmoprotectants such as glycine, betaine, choline, sarcosine, and glutamate [44]. The cryo-protectants (metabolites and proteins) suppress the aggregation of cellular proteins, stabilize phospholipid bilayers, prevent or reduce ice-crystal formation and freezing damage in bacterial cells at low temperature [45–47]. The significantly regulated water and lipid soluble compounds are mainly involved in energy conservation, carbon, protein, nucleic acid, fatty acid, and cryoprotectant biosynthesis. Energy conservation is an essential part of energy stress responses and is associated with many types of stress reactions [5,16].

In conclusion, our metabolomic study, using GC-MS and NMR, showed that the Arctic Mesorhizobium N33 regulates the levels of many compounds, and displays many molecular changes related to cold tolerance. To identify the pathways and confirm the pattern of important compounds of Arctic bacterium N33 cold adaptation ability, specific isotope dilution GC-MS or fluxomics techniques are required for accurately quantifying the metabolites in strain N33.
under cold conditions. Further investigations combining different omics technologies such as proteomics, genomics, transcriptomics, metabolomics, are required to provide a more complete system biology perspective [18] for a better understanding of the complex mechanisms of cold adaptation in the Arctic \textit{Mesorhizobium} N\textsubscript{33}.

\section*{Materials and Methods}

\subsection*{Strain Cultivation, Experimental Design and Sample Collections}

Frozen glycerol stocks of Arctic \textit{Mesorhizobium} sp. strain N\textsubscript{33} \cite{20} were used to inoculate 20 ml yeast mannitol broth (YMB) medium \cite{48} containing 200 \textmu g ml\textsuperscript{-1} streptomycin, incubated at 21°C for 5 days on a rotary shaker (180 rpm). The purity of the cells was monitored by plating on solid yeast mannitol agar (YMA) medium containing 200 \textmu g ml\textsuperscript{-1} streptomycin and 25 \textmu g ml\textsuperscript{-1} Congo red \cite{49} after 4 days of incubation at 21°C. Subsequent purity test of the strain was performed with a nodulation test on \textit{Onobrychis viciifolia} (sainfoin) \cite{48} and by 16S-rDNA sequencing \cite{50}.

A total of 8 conditions with 3 biological replicates were used in this experiment. After 3 days of growth at 21°C in YMB, 100 \mu L of pure fresh N\textsubscript{33} cells were used to inoculate 500 ml Erlenmeyer flasks containing 100 ml YMB. Flasks were incubated on a rotary shaker (180 rpm) until cells reached the mid-exponential log phase \((\text{OD}_{600} = 0.4–0.6)\), under three growing temperature conditions \((21°C = \text{control}, \ 4°C, \ 10°C; \ GT21, \ GT4 \text{ and } GT10)\). For the cold stress treatments, mid-log phase cells grown at 21°C \((T0)\) were exposed to 4°C for different times \((T1 = 2 \text{ min}, \ T2 = 4 \text{ min}, \ T3 = 8 \text{ min}, \ T4 = 60 \text{ min}, \ T5 = 240 \text{ min})\); \textbf{Figure S1} in a rotary shaker water bath \((180 \text{ rpm})\). The samples were immediately transferred into pre-chilled \((4°C)\) Falcon tubes then centrifuged \((10,000 \text{ g})\) at 4°C for 5 min. The pellets \((100 \text{ mg})\) were washed once with cold TES buffer \cite{51} to eliminate extra-cellular polysaccharides and centrifuged again. The cells were immediately quenched in liquid nitrogen \cite{52} and stored at \(-85°C\). Each 100 ml inoculated YMB provided approximately 100 mg fresh cells. In order to provide enough cell pellets, the experiments were replicated 10 times for each treatment. The pellets from 10 individual flasks per treatment were pooled together before the extraction process and considered as one biological replicate.

\subsection*{Extraction of Lipids and Water-Soluble Metabolites}

Bacterial pellets \((0.12 \text{ g dry weight})\) were transferred into a 12 ml screw capped glass vials. Lipids were extracted from the bacterial pellets according to the method of Bligh and Dyer \cite{53}. Pellets were homogenized with the Tissue-Tearor (Biospec Products, U.S.A) in 10 ml of 2:1\,(v/v) chloroform: methanol and shaken thoroughly at 150 rpm for 2 h. The extractant was centrifuged at 3000 rpm for 30 min and the supernatant was transferred into a separate vial. The residue was further extracted with 10 ml of 2:1 \,(v/v) chloroform: methanol, centrifuged again and both supernatants were pooled. Water soluble metabolites and the lipids were extracted by phase separation of a biphasic system, generated by the addition of one quarter volume of 0.88% potassium chloride solution to the chloroform: methanol extract. This mixture was shaken thoroughly for 30 min and centrifuged. The upper aqueous water-soluble metabolite layer obtained was carefully pipetted into a separate tube and purged with nitrogen

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Figure5.png}
\caption{PCA and PLS-DA of fatty acids from glycolipids present in \textit{Mesorhizobium} N\textsubscript{33} growing at constant temperatures or exposed to suboptimal 4°C. Growth temperatures: GT21 = 21°C (control); GT4 = 4°C; GT10 = 10°C. For all data, row-wise normalization was used by a pooled averaged reference samples (GT21 or T0), and data were auto scaled and log transformed. A: PCA analysis was performed on 17 fatty acids. B: PLS-DA plot showing compound changes (permutation test, \(P<0.01\)) at different times of exposure to suboptimal 4°C. T0 = 21°C (reference), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. The optimal PLS-DA model for fatty acids from glycolipids used the top three component with a \(Q^2\) = 0.75. doi:10.1371/journal.pone.0084801.g005}
\end{figure}
gas to remove traces of the solvent. The lower organic lipid layer was evaporated under nitrogen. Lipids were then resuspended in hexane and immediately used or stored under nitrogen at −20°C.

Total lipid extracts were further separated into neutral lipids, glycolipids and phospholipids classes on LC-Silica Sep Pak cartridges (Supelco) as described by Lynch and Steponkus [54]. Calculated amount of total lipid extract (10–15 mg) dissolved in 1 ml chloroform were transferred to the cartridge. Once the sample had entered the packing, residual sample was washed into the column using 2 ml chloroform followed by an additional 10 to 12 ml of the same solvent to elute neutral lipids. The glycolipids were eluted by the addition of 15 ml acetone: methanol (9:1 v/v) while phospholipids were sequentially eluted using 10 ml methanol. The fractions were dried under nitrogen, resuspended in hexane and immediately used or stored under nitrogen at −20°C.

**Preparation of Fatty Acid Methyl Esters (FAMEs)**

Fatty acid methyl esters (FAMES) were prepared according to Christie [55]. A known amount of lipids in 2:1 chloroform: methanol (v/v) was mixed with an internal standard (heptadecanoic acid C17:0, Sigma-Aldrich). The mixture was evaporated under nitrogen gas and 1 ml of the methylaing reagent (2% sulphuric acid in methanol, v/v) was added. The mixture was incubated at 80°C for 1 h, cooled on ice for 10 min after incubation and neutralized by adding 0.5 ml of a 0.5% sodium chloride solution. FAMES were extracted twice by vortexing after the addition of 2 ml aliquots of hexane. The two layers were allowed to separate and the upper hexane layer was recovered, and subjected to gas chromatographic analysis for quantification of fatty acids.

**Gas Chromatographic Analysis of FAMEs**

Gas chromatographic analysis of FAMES was done using heptadecanoic acid as the internal standard. The analysis was performed on an Agilent 6890N gas chromatography instrument coupled with an Agilent MS-5975 inert XL mass selective detector (Agilent technologies) using the electron impact (EI) ionization mode. Separation of fatty acids was achieved by injecting 2 μl of the FAMEs in a 5% phenyl 95% dimethylpolysiloxane column DB5 (Agilent J & W Scientific, 30×0.25 mm×0.25 μm). Splitless injection was performed with a constant carrier gas (helium) flow of 1 ml/min. Inlet temperature and transfer line temperatures were set at 250 and 280°C respectively. The temperature programming was as follows: an initial isotherm at 70°C was held for 1 min, raised to 76°C at a rate of 1°C/min, and from 76 to 310°C at a rate of 6.1°C/min. The MS ion source temperature was 230°C and the Quadrupole temperature was 150°C. Peak identification of fatty acids was carried out by comparison of chromatogram with mass spectral library (NIST) and against the retention times and mass spectra of the Supelco 37 component FAME mix (Sigma-Aldrich, St Louis, MO, USA).

**Figure 6. PCA and PLS-DA of fatty acids from total lipids present in Mesorhizobium N33 growing at constant temperatures or exposed to suboptimal 4°C.** Growth temperatures: GT21 = 21°C (control); GT4 = 4°C; GT10 = 10°C. For both groups of data, row-wise normalization was used by a pooled averaged reference samples (GT21 or T0), and data were auto scaled and log transformed. A: PCA analysis was performed on 13 fatty acids total. B: PLS-DA plot of total fatty acids data from GC-MS shows significant trends of the separation of compounds changes (permutation test, P<0.01) at different times of cold treatment conditions T0 = 21°C (reference), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. The optimal PLS-DA model for fatty acids from total lipids used the top one component with a Q² of 0.31. doi:10.1371/journal.pone.0084801.g006
**Figure 7.** Significant changes (P ≤ 0.05, FC ≥ 2) of fatty acids in *Mesorhizobium N₃₃* exposed to cold. GT4 = Growth at 4°C; T1 = 2 min, T2 = 4 min, T3 = 8 min, T4 = 60 min, T5 = 240 min exposure to 4°C of cells grown at 21°C; GT10 = Growth at 10°C. Observed significant changes: A: fatty acids from total lipids; B: fatty acids from neutral lipids; C: fatty acids from glycolipids and D: fatty acids from phospholipids. Arrows indicate change in concentration: green-increase, red-decrease. FC = fold changes; NA = no significant change. doi:10.1371/journal.pone.0084801.g007

| A | B | C | D |
|---|---|---|---|
| **Fatty acids total** | **Fatty acids neutral** | **Fatty acids glyco** | **Fatty acids phospho** |
| 19:1(10) | 0.068 | 19:1(10) | 0.145 | 0.026 | 19:1(10) | 0.025 | 18:2(6,9) | 0.2458 |
| 18:2(6,9) | 17.990 | 18:2(6,9) | 18:1(10) | 0.314 | 14:0 | 3.824 | 18:2(9,12) | 11.653 |
| 22:1 | 5.621 | 18:2(6,9) | 21:13 | 3.246 | 19:1(10) | 0.391 | 14:0 | 5.030 |
| 18:1(10) | 2.219 | 18:1(9) | 3.061 | 22:1(13) | 5.583 | 18:2(9,12) | 3.848 | 18:2(6,9) | 0.204 |
| 15:0 | 0.408 | 12:0 | 2.671 | 19:1(10) | 3.377 | 18:2(6,9) | 18:0 | 0.020 |
| 18:2(6,9) | 2.076 | 20:5 | 5.866 | 18:2(6,9) | 18:0 | 18:2(6,9) | 18:0 | 18:2(6,9) | 18:0 |
| **Number of fatty acids total showing significant changes** | | | |
| 20:0 | 0.269 | 16:1 | 19:0 | 9.195 | 16:0 | 0.405 | 18:1(9) | 0.138 |
| 16:0 | 0.405 | 18:1(10) | 4.068 | 14:1(11) | 45.019 | 18:0 | 0.419 | 19:1(10) | 0.458 |
| 22:1(13) | 2.780 | 16:1 | 3.236 | 18:1(9) | 3.061 | 12:0 | 2.671 | 18:1(9) | 3.061 |
| **Number of fatty acids neutral showing significant changes** | | | |
| 18:1(10) | 0.166 | 16:1(9) | 0.438 | 22:1(13) | 3.035 | 18:2(6,9) | 12.477 | 14:0 | 2.566 |
| 19:1(10) | 0.026 | 18:2(6,9) | 21:13 | 5.583 | 18:2(6,9) | 0.435 | 14:1(11) | 4.190 | 19:0 | 3.105 | 18:0 | 0.451 |
| 18:2(9,12) | 4.496 | 18:1(11) | 3.377 | 18:2(6,9) | 0.384 | 15:0 | 2.646 | 14:1(11) | 2.001 |
| **Number of fatty acids glyco showing significant changes** | | | |
| 18:1(10) | 0.026 | 18:2(9,12) | 24.583 | 18:2(9,12) | 11.653 | 14:1(11) | 44.794 | 18:0 | 0.175 |
| 18:0 | 0.204 | 18:2(6,9) | 2.495 | 14:0 | 2.269 | 19:1(10) | 0.178 |
| 16:1(7) | 3.848 | 18:2(6,9) | 0.434 | 18:2(6,9) | 0.285 | 16:0 | 0.499 |
| 14:0 | 3.943 | 18:2(6,9) | 10.467 | 18:2(6,9) | 6.956 | 18:2(6,9) | 16.624 |

**Legend:**
- **GT4:** Growth at 4°C
- **T1:** 2 min, **T2:** 4 min, **T3:** 8 min, **T4:** 60 min, **T5:** 240 min exposure to 4°C of cells grown at 21°C
- **GT10:** Growth at 10°C
- **FC:** Fold changes
- **NA:** No significant change
GC-MS Method for Identification of Water-Soluble Metabolites

To derivatize the metabolites for GC-MS analysis, 40 μL of methoxamine hydrochloride (Sigma-Aldrich) in pyridine was added to the water-soluble extracts and incubated at room temperature for 16 hours. Then 50 μL (N-Methyl-N-trifluoroaceticamide) MSTFA with 1% TMCS (Trimethylchlorosilane) derivatization agent were added and incubated at 37°C for 60 minutes on a hotplate. The samples were vortexed twice throughout the incubation period to ensure complete dissolution. Samples were refrigerated at 4°C for no longer than 48 hours before analysis in order to avoid any degradation of the derivatized compounds.

Derivatized extracts were analyzed using an Agilent 7890-5975C GC-MS instrument operating in an Electron Impact (EI) ionization mode. For GC-MS analysis, 2 μL of the derivatized samples were injected using a split/splitless injector with a split ratio of 5:1 onto a HP-5MS capillary column (30 m × 250 μm × 0.25 μm). The helium carrier gas was set to a flow rate of 1 mL/min and the initial oven temperature was set to 70°C. The temperature was increased at 1°C/min to 76°C, and then at 6.1°C/min to 310°C. The total run time was 45 minutes. The full scan mode of the quadrupole MS was used at a mass range of 50–500 m/z, with a solvent delay of 6 minutes. The ion source temperature was 230°C and the Quadrupole temperature was 150°C. In GC-MS, a faster scan speed generally provides more data points across a chromatographic peak, but it tends to lower the ion statistics. In contrast, a slower scan rate produces few scans over the peak and results in better spectra. The scan speed of our quadrupole MS was optimized over a number of experiments (~42.9 mg/ml extracted cells) was dissolved in 500 μL of 50 mM NaH2PO4 buffer pH 7. Thirty five μL of D2O and 15 μL of a buffer solution (0.5 mM DSS (disodium-2,2-dimethyl-2-silapentane-5-sulphonate) and 0.47% NaN3) were added to the sample. The sample amount in the final assay volume (350 μL) was 15 mg. The sample solution was vortexed for 1 minute, sonicated for 30 minutes, and transferred to a standard Shigemi microcell NMR tube for subsequent spectral analysis.

NMR Spectroscopy

All 1H-NMR spectra were collected on a 500 MHz Inova (Varian Inc., Palo Alto, CA) spectrometer equipped with a 5 mm HCN Z-gradient pulsed-field gradient (PFG) room-temperature probe. 1H-NMR spectra were acquired at 25°C using the first transient of the NOESY-presaturation pulse sequence, which was chosen for its high degree of quantitave accuracy. Spectra were collected with 256 transients using a 4s acquisition time and a 1s recycle delay.

NMR Compound Identification and Quantification

All FIDs (free induction decays) were zero-filled to 64k data points and subjected to line broadening of 0.5 Hz. The singlet produced by the DSS methyl groups was used as an internal standard for chemical shift referencing (set to 0 ppm) and for quantification. All 1H-NMR spectra were processed and analyzed using the Chenomx NMR Suite Professional software package version 6.0 (Chenomx Inc., Edmonton, AB). The Chenomx NMR Suite software allows for qualitative and quantitative analysis of an NMR spectrum by manually fitting spectral signatures from an internal database of reference spectra to the full NMR spectrum [56]. Specifically, the spectral fitting for each metabolite was done using the standard Chenomx 500 MHz metabolite library. Typically 90% of all visible peaks were assigned to a compound.
and more than 90% of the spectral area could be routinely fit using the Chenomx spectral analysis software. Most of the visible peaks are annotated with a compound name. It has been previously shown that this fitting procedure provides absolute concentration accuracies of 90% or better. Each spectrum was processed and analyzed by at least two NMR spectroscopists to minimize compound misidentification and misquantification. We used sample spiking to confirm the identities of assigned compound. The confirmations of the spiking with standard and original peaks in experiments for those unexpected metabolites (i.e. phenylacetate and N-carbamoyl-beta-alanine) and for those hard to assign (i.e. malonic acid and oxypurinol) are shown in Figure S12 (A–D). Sample spiking involves the addition of 20–200 μM of the suspected compound to selected samples and testing whether the relative NMR signal intensity changed as expected.

Data Processing for Statistical Analysis
To visualize the compounds, and compare the metabolite changes (composition and concentrations) of all cold perturbation treatments with the control (N\textsubscript{33}, cells grown at 21°C = GT21 or T0), several multivariate and univariate analytical methods were applied. The water and lipid soluble compound concentration data tables were arranged with samples in column and compounds in rows. Data tables were formatted as comma separated values (.csv). Data tables with three growing temperatures (GT21, GT4, GT10) and 6 time points (T0, T1, T2, T3, T4, T5) were uploaded to the MetaboAnalyst 2.0 server [http://www.metaboanalyst.ca] [57] and analysed separately. In both groups of data metabolite, data were unpaired and analysed using multivariate and univariate methods. To reduce any possible variance and to improve the performance for downstream statistical analysis, metabolites data generated by GC-MS and NMR were normalized using MetaboAnalyst’s normalization protocols [57]. For multivariate analysis (PCA and PLS-DA), Row-wise normalization was performed for all metabolite data by comparing the samples with a pooled average of reference samples (cells grown at 21°C) to make each sample comparable to one another. To make the metabolite concentration values more comparable among different compounds, several different types of column-wise normalization were performed. The fractions of fatty acids (from total, neutral, phospho-, and glyco- lipids) were log transformed and analysed individually. The fatty acids data were also normalized using auto scaling (mean-centered and divided by the standard deviation of each variable). Same normalization procedures were performed for NMR data. Univariate analysis was applied to calculate the fold change (FC), volcano plots, and statistical significance (t-test and one way ANOVA) assessed. Since FC is calculated as the ratio between two group means (sample over control), the column-wise normalization (i.e. log transformation/mean-centering) will significantly change the absolute values, thus in univariate analysis data were used before column normalization [58]. To assess the degree of metabolite concentration changes, heat maps and hierarchical clustering were performed using the MetaboAnalyst 2.0 software [57]. Heatmaps were created based on the Pearson distance measure and the Ward clustering algorithm, displayed for top 25 features selected by analysis of variance (ANOVA) using a significance level of P≤0.05, and post-hoc analysis of Fisher’s LSD.

Supporting Information
Figure S1 Experimental plan used to study metabolomics of cold adaptation of the Arctic Mesorhizobium N\textsubscript{33}. Bacteria were cultivated in yeast manitol broth (YMB) medium at a constant temperature of 21°C (GT21), 4°C (GT4) or 10°C (GT10). For the effect of time of exposure to cold, N\textsubscript{33} cells were grown at 21°C (T0) and then exposed to 4°C for: 2 min (T1), 4 min (T2), 8 min (T3), 60 min (T4) and 240 min (T5). Each treatment is color coded, and the color codes are used in all figures of this manuscript.

Figure S2 Heatmap visualization of water soluble metabolites present in Mesorhizobium N\textsubscript{33} during growth at different temperatures. GT21 = growth at 21°C (control); GT4 = growth at 4°C; GT10 = growth at 10°C. Data were row-wise normalized by a pooled averaged reference sample (GT21), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 29 water soluble metabolites and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows for each growth temperature used a distinct effect Conditions GT21 and GT10 represent close change trends of the water soluble metabolites. However some compounds have shown slightly different levels of accumulations. Metabolites of the cells grown at 4°C (GT4) are clustered in a distinct group far from those of GT10 and GT21. Most influential compounds that were highly accumulated during constant growth at 4°C include isobutyrate, sarcosine, threonine, and valine.

(TIF)

Figure S3 Heatmap visualization of water soluble metabolites present in Mesorhizobium N\textsubscript{33} exposed to suboptimal 4°C for various times. T0 = 21°C (control), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. Data were row-wise normalized by a pooled averaged reference sample (T0), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 29 water soluble metabolites and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of metabolite changes under each time of exposure to low temperature.

(TIF)

Figure S4 Heatmap visualization of fatty acids from neutral lipids present in Mesorhizobium N\textsubscript{33} during growth at different temperatures. GT21 = growth at 21°C (control); GT4 = growth at 4°C; GT10 = growth at 10°C. Data were row-wise normalized by a pooled averaged reference samples (GT21), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 16 fatty acids from neutral lipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The fatty acids were clustered in 5 groups. Conditions GT21 and GT10 represent a close change trends in fatty acids. Metabolites of cells grown at the 4°C (GT4) are clustered in distinct group and far from those of GT10 and GT21. 

(TIF)

Figure S5 Heatmap visualization of fatty acids from neutral lipids present in Mesorhizobium N\textsubscript{33} exposed to suboptimal 4°C for various times. T0 = 21°C (control), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. Data were row-wise normalized by a pooled averaged reference sample (T0), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 17 fatty acids from neutral lipids and is shown at the top and side of the panel. Brown
Figure S6 Heatmap visualization of fatty acids from phospholipids present in *Mesorhizobium N33* during growth at different temperatures. GT21 = growth at 21°C (control); GT4 = growth at 4°C; GT10 = growth at 10°C. Data were row-wise normalized by a pooled averaged reference samples (GT21), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 17 fatty acids from phospholipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of the fatty acids changes Metabolites of the cells grown at 21°C (GT21) are clustered in distinct group and far from those GT10 and GT4.

(TIF)

Figure S7 Heatmap visualization of fatty acids from phospholipids present in *Mesorhizobium N33* exposed to suboptimal 4°C for various times. T0 = 21°C (control), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. Data were row-wise normalized by a pooled averaged reference sample (T0), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 17 fatty acids from phospholipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of metabolic changes under each time of exposure to low temperature.

(TIF)

Figure S8 Heatmap visualization of fatty acids from glycolipids present in *Mesorhizobium N33* during growth at different temperatures. GT21 = growth at 21°C (control); GT4 = growth at 4°C; GT10 = growth at 10°C. Data were row-wise normalized by a pooled averaged reference samples (GT21), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 17 fatty acids from glycolipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of the fatty acids changes under each time of exposure to low temperature.

(TIF)

Figure S9 Heatmap visualization of fatty acids from glycolipids present in *Mesorhizobium N33* exposed to suboptimal 4°C for various times. T0 = 21°C (control), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. Data were row-wise normalized by a pooled averaged reference sample (T0), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 17 fatty acids from glycolipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of metabolic changes under each time of exposure to low temperature.

(TIF)

Figure S10 Heatmap visualization of fatty acids from total lipids present in *Mesorhizobium N33* during growth at different temperatures. GT21 = growth at 21°C (control); GT4 = growth at 4°C; GT10 = growth at 10°C. Data were row-wise normalized by a pooled averaged reference samples (GT21), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 13 fatty acids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows distinct trends of the metabolite changes at 21°C (GT21), 10°C (GT10) and 4°C (GT4). The fatty acids were grouped in 3 main clusters and 8 sub-clusters in Heatmaps. Conditions GT4 represents distinct trends of metabolite changes compared to the metabolites of the cells grown at 21°C and 10°C. Out of 13 fatty acids of total lipids, at least 6 fatty acids showed accumulation at 4°C and 7 compounds showed down regulations.

(TIF)

Figure S11 Heatmap visualization of fatty acids from total lipids present in *Mesorhizobium N33* exposed to suboptimal 4°C for various times. T0 = 21°C (control), T1 = 2 min; T2 = 4 min; T3 = 8 min; T4 = 60 min; T5 = 240 min exposure to 4°C of cells grown at 21°C. Data were row-wise normalized by a pooled averaged reference sample (T0), and were auto scaled and log transformed. Hierarchical clustering was performed based on Pearson’s distance on 13 fatty acids from total lipids and is shown at the top and side of the panel. Brown and blue colors represent an increase and decrease of a metabolite. The heatmap visualization shows different trends of metabolic changes under each time exposure at low temperature.

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Figure S12 Confirmations by spiking with standard and original peaks of unexpected metabolites. A) Malonic acid, B) Oxypurinol, C) Phenylacetate, D) N-Carbamoyl-beta-alanine.

(TIF)

Table S1 Low temperature effects on the concentration (μM) of 29 water-soluble metabolites determined by NMR in Arctic *Mesorhizobium N33*.

(DOCX)

Table S2 Low temperature effects on the fatty acid composition of total lipids determined by GC-MS in Arctic *Mesorhizobium N33* (expressed as mole % of total fatty acids).

(DOCX)

Table S3 Low temperature effects on the fatty acid composition of neutral lipids determined by GC-MS in arctic *Mesorhizobium strain N33* (expressed as mole % of total neutral fatty acids).

(DOCX)

Table S4 Low temperature effects on the fatty acid composition of glycolipids determined by GC-MS in arctic *Mesorhizobium strain N33* (expressed as mole % of total glycolipids).

(DOCX)

Table S5 Low temperature effects on the fatty acid composition of phospholipids determined by GC-MS in arctic *Mesorhizobium strain N33* (expressed as mole % of total phospholipids).

(DOCX)
Author Contributions
Conceived and designed the experiments: AG SL. Performed the experiments: AG RK. Analyzed the data: AG. Contributed reagents/

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