VIB1, a Link between Glucose Signaling and Carbon Catabolite Repression, Is Essential for Plant Cell Wall Degradation by Neurospora crassa

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Abstract

Filamentous fungi that thrive on plant biomass are the major producers of hydrolytic enzymes used to decompose lignocellulose for biofuel production. Although induction of cellulases is regulated at the transcriptional level, how filamentous fungi sense and signal carbon-limited conditions to coordinate cell metabolism and regulate cellulolytic enzyme production is not well characterized. By screening a transcription factor deletion set in the filamentous fungus Neurospora crassa for mutants unable to grow on cellulolytic materials, we identified a role for the transcription factor, VIB1, as essential for cellulose utilization. VIB1 does not directly regulate hydrolytic enzyme gene expression or function in cellulolytic inducer signaling/processing, but affects the expression level of an essential regulator of hydrolytic enzyme genes, CLR1 and CLR2. Transcriptional profiling of a Δvib-1 mutant suggests that it has an improper expression of genes functioning in metabolism and energy and a deregulation of carbon catabolite repression (CCR). By characterizing new genes, we demonstrate that the transcription factor, COL26, is critical for intracellular glucose sensing/metabolism and plays a role in CCR by negatively regulating cre-1 expression. Deletion of the major player in CCR, cre-1, or a deletion of col-26, did not rescue the growth of Δvib-1 on cellulose. However, the synergistic effect of the Δcre-1; Δcol-26 mutations circumvented the requirement of VIB1 for cellulase gene expression, enzyme secretion and cellulose deconstruction. Our findings support a function of VIB1 in repressing both glucose signaling and CCR under carbon-limited conditions, thus enabling a proper cellular response for plant biomass deconstruction and utilization.

Introduction

Bioconversion of lignocellulose biomass to simple sugars holds great promise in next-generation biofuel production and relies on a complex repertoire of proteins for enzymatic deconstruction of plant cell walls [1]. Many filamentous fungi have evolved to utilize cellulolytic materials and are capable of producing a wide spectrum of enzymes, but only a few species have been harnessed for industrial usage [2]. Further improvement in fungal cellulolytic enzyme production is desired to make biofuel production cost-competitive, but this relies on a better understanding of the molecular basis of networks involved in carbon sensing and regulatory aspects associated with induction of gene expression of hydrolytic enzymes [3].

Cellulolytic enzyme production and secretion is a unique attribute of filamentous fungi, and efforts to identify important factors in enzyme production led to the discovery of a number of transcriptional activators and repressors. For example, the transcription factor XlnR/XYR1 positively regulates expression of cellulase and hemicellulase genes in Aspergillus niger and Trichoderma reesi, respectively [4–7]. In Neurospora crassa, the transcription factors CLR1 and CLR2 are essential for growth on cellulose and are required for expression of a ~212 gene regulon that is induced in response to cellohextrins, such as cellobiose [8,9] (Figure 1). In Aspergillus nidulans and A. oryzae, a clb-2 homolog, called clbB/manR, respectively, is also essential for cellulase gene expression and activity [8,10,11]. Additional transcriptional regulators that promote expression of some genes encoding hydrolytic enzymes have also been identified, including menA in A. nidulans [12], clbR in A. aculeatus [13], and aceI and bgIR in T. reesi [14,15].

In addition to induction, cellulase gene expression is also subject to carbon catabolite repression (CCR), which functions when a favorable carbon source, such as glucose, is present [3,16,17]. The most well-characterized transcription factor involved in CCR in filamentous fungi is CreA/CRE1. Deletion of creA/cre-1 alleviates some aspects of CCR for cellulolytic enzyme expression in Aspergilli [18–22], T. reesei [23–25], Penicillium decumbens [2] and N. crassa [26,27]. In A. nidulans, repression by CreA occurs both by binding to promoters of hemi-cellulase genes as well as repressing expression of transcriptional activators [28]. Other factors including creB/cre2, creC, creD, lim1, and aceI were also reported to promote CCR in different fungal species via unknown mechanisms [29–37]. The strength of CCR is tuned by glucose sensing and signaling, although crosstalk between these two regulatory systems is not well understood. In N. crassa, RCO3,
Author Summary

Many filamentous fungi that grow on plant biomass are capable of producing lignocellulose enzymes to break down plant cell walls into utilizable sugars, thus holding great potential in reducing the cost of the next-generation biofuels. Cellulase production is subject to induction by the presence of plant biomass components and to repression by the availability of easily metabolized sugars, such as glucose. Genes required for repression of cellulase gene expression when preferred carbon sources are present (carbon catabolite repression) and those that play a role in mediating glucose sensing/metabolism have been identified in filamentous fungi, but the mechanisms involved in crosstalk between repression versus induction of cellulase gene expression is poorly understood. Here, we report the identification and functional characterization of VIB1, a transcription factor essential for plant cell wall deconstruction in Neurospora crassa and COL26, a transcription factor that functions in glucose sensing/metabolism and regulation of CCR. We show that disabling CRE1 repression and modulating the glucose response by deletion of col-26 restored growth of the Δvib-1 mutant on cellulose. Our findings are particularly important in understanding the molecular basis of enzyme production that could allow a further strain improvement for plant biomass deconstruction.

Figure 1. Cellulase production in N. crassa is regulated by cellobiose induction and CCR. CCR is decreased in absence of cellobiose, allowing scouting enzymes to liberate cellobiose from cellulose. Cellobiose (or a derivative) results in activation of the transcription factor CLR2, which induces expression of transporters for cellobiose. CLR1 and CLR2 drive cellulase gene expression. Both intracellular and extracellular β-glucosidases enzymes catalyze conversion of cellobiose to glucose, which can trigger carbon catabolite repression via glucose sensing mechanisms and transcriptional repression by CRE1.

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Results

Deletion of vib-1 causes a growth defect on cellulose biomass

Screening of a transcription factor deletion set of N. crassa strains [46] for ability to deconstruct crystalline cellulose showed that a strain carrying a deletion of the vib-1 gene (FGSC11309) failed to grow on Avicel (Figure 2A). Since functional vib-1 is required for extracellular protease secretion in response to carbon and nitrogen starvation in N. crassa [48,49], we hypothesized that the Δvib-1 mutant might be unable to respond to complex extracellular carbon sources. In support of this hypothesis, the Δvib-1 mutant also exhibited slow growth on xylan. Growth defects were accompanied by barely detectable extracellular enzyme activity towards crystalline cellulose and low extracellular xylanase activity (Figure 2B and S1A). In contrast, the Δvib-1 mutant accumulated a similar amount of mycelial biomass as the WT strain when inoculated into minimal media containing simple sugars (sucrose, cellobiose or xylose) (Figure S1B). The introduction of an ectopic copy of vib-1 (Pvib-1) completely restored the growth defects on Avicel of the Δvib-1 mutant, as well as the secretome and cellulolytic enzyme activity of culture supernatants (Figure 2B).

To test the hypothesis that the role of VIB1 in cellulose utilization is conserved in other filamentous fungi, especially in fungi used in industrial production of cellulolytic enzymes, we carried out complementation tests using the vib-1 ortholog from T. reesei (EGR52133; Trvib1); TrVIB1 and N. crassa Vib1 share 49% amino acid identity. Constitutive expression of Trvib1 in a N. crassa Δvib-1 mutant fully restored the growth and cellulolytic enzyme activity (Figure 2B). The Trvib1 strain also recapitulated most of the secretome of N. crassa WT and Pvb-1 strains on Avicel (Figure 2C). These results suggest that vib-1 is functionally conserved for the utilization of cellulose in filamentous ascomycete fungi.

The Δvib-1 mutant shows an inappropriate temporal and spatial conidiation pattern. These phenotypes are correlated with differential localization of VIB1-GFP in vegetative hyphae versus conidiophores [48]. As conidiation is regulated by glucose
Figure 2. Deletion of vib-1 abolishes production of cellulases and utilization of cellulosic material. (A) Growth of WT and Δvib-1 on Avicel after 4 days; growth of WT is indicated by formation of orange mycelia, versus no growth of the Δvib-1 mutant. (B) Cellulase activity from 4-day old culture supernatants from Avicel-grown cultures of WT, the Δvib-1 mutant, the Pviib-1 strain (constitutive expression of vib-1 in a Δvib-1 strain) and the PTrvib-1 strain (constitutive expression of T. reesei vib1 in a Δvib-1 strain). Cellulase activity was measured using Avicel as a substrate and represented by the amount of glucose and cellobiose released. The equivalent of glucose from cellobiose was calculated and represented by the light gray bar. (C) The secretomes of strains analyzed in panel B are shown.

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limitation [50], we assessed whether differential localization of VIB1 was also associated with cellulose utilization. We examined a strain in which we replaced the resident vib-1 gene with a functional vib-1-gfp construct. Nuclear localization of VIB1-GFP was observed in hyphae and localization was independent of carbon source, either following a shift to Avicel for 2 hrs (Figure S1C) or after prolonged growth.

Constitutive expression of clr-2 restores cellulose utilization in the Δvib-1 mutant

Previous comparative RNA-seq analysis of WT revealed that 212 genes are significantly differentially expressed under cellulosic conditions, a gene set referred to as the “Avicel regulon” [8]. To determine whether the defect in cellulase secretion and activity in the Δvib-1 mutant was due to failure to induce cellulase gene expression versus a defect in cellulase secretion, we assessed genome-wide expression differences via RNA-seq between the WT and Δvib-1 strains following a shift for 4 hrs from sucrose medium to either carbon-free or Avicel medium. Of the 212 genes in the Avicel regulon, 91 genes were expressed at a significantly lower level in the Δvib-1 mutant versus WT under Avicel conditions (cutoff: Padj <0.05 and fold change ≥2; Table S1). This gene set includes the essential cellulose transcription factor gene, clr-2 and 43 carbohydrate-active-enzymes (CAZy) from 27 different families (Carbohydrate Active Enzymes database: http://www.cazy.org/) [51].

CLR1 and CLR2 are strictly required for full expression of 140 genes within the Avicel regulon [8,10]; 62 of these genes were identified in the 91-gene set that showed low expression in the Δvib-1 mutant. Although expression of clr-1 was not significantly different from WT in the Δvib-1 mutant, the expression of clr-2 was significantly reduced (FPKM: 107±43 in WT; 66±27 in Δvib-1 for clr-1 versus 171±10 in WT; 39±14 in Δvib-1 for clr-2). Importantly, constitutive expression of clr-2 (Pc clr-2) in minimal medium without cellulosic inducers recapitulates the response of N. crassa to crystalline cellulose, including the secretion of active cellulolytic enzymes [10]. The reduced transcription of clr-2 in the Δvib-1 mutant (Table S1) suggested that constitutive expression of clr-2 might suppress the cellulose utilization defect in the Δvib-1 mutant. To test this hypothesis, we constructed a Pc clr-2; Δvib-1 strain and evaluated its ability to secrete cellulases and utilize Avicel in comparison to the Pc clr-2, the Δvib-1 and WT strains. In support of our hypothesis, the Pc clr-2; Δvib-1 strain showed restoration of protein secretion and cellulolytic activity to near WT levels (Figure 3A). Although the clr-2 expression levels in the Pc clr-2 strain were at a similar level to a WT strain after a 4 hr shift to Avicel (Figure S4), the Pc clr-2; Δvib-1 mutant showed a ∼3.5 fold increase in clr-2 expression level under the same conditions.

To evaluate the functions of VIB1 versus CLR2 in regulating cellulose utilization, we generated RNA-seq data from the Pc clr-2; Δvib-1 mutant that was shifted for 4 hrs from sucrose medium to carbon-free medium and compared it to previously obtained data for Pc clr-2 [10]. Analysis of genes encoding different CAZy family proteins revealed a similar pattern of expression between the Pc clr-2 and the Pc clr-2; Δvib-1 strains (Figure 3B), consistent with our hypothesis that VIB1 functions upstream of clr-2 in response to cellulose. To differentiate whether any Avicel-regulon genes that showed decreased expression levels in Δvib-1 mutant were due to the vib-1 deletion rather than a low level of clr-2 expression, we performed hierarchical clustering of expression patterns of the Avicel-regulon genes across 6 RNA-seq experiments (WT and Δvib-1 shifted to no carbon or Avicel and the Pc clr-2 and the Pc clr-2; Δvib-1 strains shifted to no carbon); 4 major expression groups were identified (Figure 3C and Table S1). Two groups (group 1 and 3) were CLR2-regulon genes that are vib-1 independent. Group 1 consisted of 54 genes whose
expression was fully induced by constitutive expression of \textit{clr-2} regardless of the presence or absence of \textit{vib-1}, including 33 of the 43 CAZy proteins (Table S1). Group 3 consisted of 8 genes whose expression was partially induced by \textit{clr-2}, but still in a \textit{vib-1}-independent manner. The fourth group of genes included 14 \textit{vib-1} modulated genes. These genes were partially induced in \textit{Δvib-1} on Avicel, but remained repressed in both the \textit{Pc clr-2} and the \textit{Pc cl-2}; \textit{Δvib-1} strains under no carbon conditions (Table S1). Expression of these genes is likely induced by the cellulolytic cascade pathways upstream of CLR2 or other components present.

**Figure 3. Constitutive expression of \textit{clr-2} rescued the cellulase production defect of the \textit{Δvib-1} mutant.** (A) Protein concentration and cellulase activity in a \textit{Δvib-1} mutant versus a \textit{Δvib-1} strain constitutively expressing \textit{clr-2} (\textit{Pc clr-2}; \textit{Δvib-1}) and WT and a \textit{Pc cl-2} strain under Avicel conditions. (B) Expression levels from RNA-seq data of genes encoding major classes of CAZy proteins from WT and \textit{Δvib-1} shifted to Avicel versus the \textit{Pc cl-2} and \textit{Pc cl-2}; \textit{Δvib-1} strains shifted to minimal media with no carbon source. FPKM (Fragment Per KiloBase per exon per Megabase mapped) for individual genes were averaged between three biological replicates and pooled by CAZy class. (C) Hierarchical clustering of FPKM for 91 Avicel-regulon genes in the \textit{Δvib-1} mutant and WT on Avicel (Av) and the \textit{Δvib-1}, WT, \textit{Pc cl-2} and the \textit{Pc cl-2}; \textit{Δvib-1} strains switched to no carbon conditions (Nc). Results are displayed as heat maps with log (FPKM) from minimum (bright blue) to maximum (bright yellow). doi:10.1371/journal.pgen.1004500.g003
in commercial Avicel preparations, such as a low concentration of hemicellulose [9]. The second group consisted of 15 genes that were induced by constitutive \( clr-2 \) expression under no carbon conditions but in \( vib-1 \)-dependent manner. This gene set included a pectate lyase (NCU06326), a BNR/Asp-box repeat protein predicted to have exo-\( \beta \)-L-1,3-arabinamidase activity (NCU09924), a \( \beta \)-xylosidase (NCU09923/gh3-7), an extracellular \( \beta \)-1,4-D-glucosidase (NCU04952/gh3-4), a \( \beta \)-1,3-glucosidase (NCU09904), a starch binding domain-containing protein (NCU08746), a LysM domain-containing protein (NCU05319), a putative methyltransferase (NCU05501), and 6 hypothetical proteins. Six genes in this set encode proteins predicted to enter the secretory pathway (Table S1).

**VIB1 functions upstream of the inducer signal**

Our epistasis experiments indicated that \( vib-1 \) functions upstream of \( clr-2 \), suggesting that VIB1 could be involved in signal molecule processing that leads to CLR1 activation and thus \( clr-2 \) expression (Figure 1). In *N. crassa*, a strain carrying deletions of genes encoding two extracellular \( \beta \)-glucosidases and an intracellular \( \beta \)-glucosidase (\( \Delta3\beta G \)), recapitulates the cellulolytic response when the \( \Delta3\beta G \) strain is exposed to cellobiose [9]. These data indicate that cellobiose (or a derivative) functions as a cellulose signal that results in the induction of cellulolytic genes and subsequent secretion of cellulase enzymes. This cellobiose-induced cellulase gene expression and secretion is dependent upon functional \( clr-2 \) gene, as the \( \Delta3\beta G; \Delta clr-2 \) mutant is unable to produce cellulolytic enzymes in response to Avicel or cellobiose (unpublished data). We therefore asked if VIB1 plays a role in induction via signal processing. To test this hypothesis, we created a \( \Delta3\beta G; \Delta vib-1 \) quadruple mutant and asked whether the \( \Delta3\beta G; \Delta vib-1 \) mutant could induce cellulase gene expression in response to cellobiose. Following a switch from sucrose to either carbon, or 0.2% cellobiose, or 2% Avicel for 4 hrs, the induction of two major cellulase genes, \( cbb-1/NCU07340 \) and \( gh5-1/NCU00762 \) were significantly induced in the \( \Delta3\beta G; \Delta vib-1 \) and the \( \Delta3\beta G \) strains, but not in the \( \Delta vib-1 \) strain (\( p<0.05 \)) (Figure 4A).

The restoration of cellulase gene expression in the \( \Delta3\beta G; \Delta vib-1 \) strain when exposed to cellobiose was accompanied by enzyme production and activity. Similar to the \( \Delta3\beta G \) mutant, the \( \Delta3\beta G; \Delta vib-1 \) strain accumulated biomass more slowly on cellobiose than WT or the \( \Delta vib-1 \) mutant due to the slow conversion of cellobiose to glucose (0.51±0.11 g/L, and 0.63±0.0 g/L for the \( \Delta3\beta G \) and the \( \Delta3\beta G; \Delta vib-1 \) strains, respectively, versus 3.83±0.19 g/L and 3.62±0.11 g/L for WT and \( \Delta vib-1 \), respectively). However, despite less biomass accumulation, both the \( \Delta3\beta G \) and the \( \Delta3\beta G; \Delta vib-1 \) strains showed significantly more enzyme activity than WT and the \( \Delta vib-1 \) strains on 2% cellobiose (Figure 4B). When grown on medium containing 2% Avicel as a sole carbon source, the \( \Delta3\beta G \) and \( \Delta vib-1 \) strain showed significantly higher enzyme activity than WT and the \( \Delta vib-1 \) strains on 2% cellobiose (Figure 4B). When grown on medium containing 2% Avicel as a sole carbon source, the \( \Delta3\beta G \) and \( \Delta vib-1 \) strain showed significantly higher enzyme activity than WT and the \( \Delta vib-1 \) strains on 2% cellobiose (Figure 4B).

**Comparative analysis of transcriptomes of WT and the \( \Delta vib-1 \) strain**

In addition to induction, cellulolytic enzyme production requires proper nutrient sensing and relief from carbon catabolite repression (CCR) (reviewed in [17,52]). We therefore hypothesized that the \( \Delta vib-1 \) mutant might be defective in either nutrient sensing and/or relieving CCR in response to Avicel. To test this hypothesis, we first compared RNA-seq data of the \( \Delta vib-1 \) mutant when shifted from sucrose to carbon-free media versus a shift from sucrose to Avicel media. This comparison revealed 770 differentially expressed genes (cutoff: \( \text{Padj}<0.01 \) and fold change >2) (Table S2). We then compared these genes were expressed in WT under no carbon versus Avicel conditions using a previously published RNA-seq dataset [8]. Hierarchical clustering analysis of expression patterns of these 770 genes revealed three gene clusters (Figure 5) (Table S2).
The first cluster contained 237 genes whose expression pattern was similar between the \( \Delta \text{vib-1} \) and WT strains. This gene set was expressed at low levels under no carbon conditions but induced to higher levels upon exposure to Avicel. This group contained 51 CAZy proteins, \( clr-1 \) and \( clr-2 \), all three cellodextrin transporters (\( cdt-1 \), \( cdt-2 \), and \( cbt-1 \)) \cite{44,53,54} and 102 hypothetical proteins. This gene set overlapped the WT Avicel-regulon for 143 genes, suggesting that cellulosic induction still occurred in the \( \Delta \text{vib-1} \) mutant albeit at a low level.

The second cluster consisted of 173 genes whose expression pattern was also similar between WT and the \( \Delta \text{vib-1} \) strain. However, in contrast to the first gene set, the expression level of these 173 genes was higher under carbon-free conditions. This set included 7 CAZy proteins, three conidiation-specific proteins (\( NCU08769/\text{con-6} \), \( NCU07325/\text{con-10} \), \( NCU09235/\text{con-8} \)), a high affinity glucose transporter/\( NCU08152 \), and 103 hypothetical proteins. Genes in this cluster may encode proteins that function in a general response to carbon starvation.

The third cluster consisted of 360 genes whose expression pattern between no carbon and Avicel conditions was different in the \( \Delta \text{vib-1} \) mutant as compared to the WT strain. This gene set showed consistently higher expression in the \( \Delta \text{vib-1} \) mutant on Avicel medium as compared to carbon-free medium (Figure 5). Only 7 genes encoding CAZy proteins were in this set and 169 genes were annotated as hypothetical. An enrichment in the categories of metabolism and energy, particularly, degradation of glycine (\( p = 2.37 \times 10^{-03} \)), nitrogen, sulfur and selenium metabolism (\( p = 8.00 \times 10^{-03} \)), purine nucleotide/nucleoside/nucleobase catabolism (\( p = 2.49 \times 10^{-05} \)), isoprenoid metabolism (\( p = 8.63 \times 10^{-04} \)), respiration (\( p = 3.34 \times 10^{-04} \)), metal binding (\( p = 6.18 \times 10^{-04} \)), and mitochondrial transport (\( p = 2.94 \times 10^{-03} \)) was observed. These data suggested that the \( \Delta \text{vib-1} \) mutant was improperly responding to carbon-limited conditions as compared to a WT strain.

Within the gene set that showed increased expression level in the \( \Delta \text{vib-1} \) mutant on Avicel were genes involved in CCR. This gene set included \( cre-1/NCU08807 \), \( creD/NCU03887 \), \( creB/NCU08378 \) and \( bgIR/NCU07788 \) (Table S2). Although the role of \( cre-1 \) in CCR and cellulose utilization is established in \( N. \ crassa \) \cite{26,27}, the function of the \( creB \) and \( creD \) homologs in cellulolytic enzyme production were uncharacterized. In \( N. \ crassa \),
NCU07788/BglR was previously characterized in a transcription factor deletion screen and was named *col-26* for its colonial phenotype on minimal sucrose medium [46].

### The identification and characterization of new proteins involved in carbon sensing

To determine whether homologs of the CCR genes that showed increased expression in the *Avib*-1 mutant play a role in cellulose deconstruction, we first measured protein concentration and cellulase enzyme activity in supernatants from the Δ*col-26*, ΔNCU08378/creB, and ΔNCU03887/creD mutants grown on Avicel for 7 days: none of the mutants showed significantly different cellulase activity than WT (Figure S3). To test if these genes are involved in CCR, we evaluated resistance of WT and the mutants to 2-deoxy-glucose (2-DG). The compound 2-DG is an analogue of glucose that cannot be metabolized and is often used to select for, or evaluate, impairment of CCR and glucose repression in filamentous fungi [39,55–57]. In strains with functional CCR, 2-DG is phosphorylated, thus activating CCR, resulting in the inability of the strain to grow on alternative carbon sources; strains with impaired CCR are insensitive to 2-DG exposure. When 2% cellobiose and 0.2% 2-DG were used as carbon sources, only the Δ*cre*-1 and the Δ*col-26* mutants showed 2-DG resistance, which was more obvious when Avicel instead of cellobiose was used as a carbon source (Figure 6A). These data implicated COL26 in CCR in *N. crassa*.

To confirm the role of COL26 in CCR, we tested CCR functionality using allyl alcohol (AA). As reported for *M. oryzae* [41], when CCR is impaired, alcohol dehydrogenase is expressed and will convert AA into toxic acrylaldehyde. Thus, strains with impaired CCR exhibit AA sensitivity, while strains with functional CCR are insensitive. As predicted, the Δ*cre*-1 mutant was sensitive to AA, but the Δ*col-26* mutant, similar to WT, was insensitive (Figure 6B). These data indicated that CCR was still functional in the Δ*col-26* mutant. To reconcile the different results for the Δ*col-26* mutant with respect to CCR, we analyzed growth of the Δ*cre*-1 and the Δ*col-26* mutants on different simple carbon sources. When grown on MM media with 2% glucose, fructose, sucrose, or cellobiose as the sole carbon source, the Δ*cre*-1 mutant accumulated a similar amount of biomass to the WT strain (Figure 7A). However, the Δ*col-26* mutant exhibited a severe growth defect on glucose, fructose and sucrose, consistent with its colonial designation [46], but only a moderate growth defect on cellobiose (Figure 7A).

The fact that the Δ*col-26* mutant grew much better on cellobiose as compared to glucose, fructose, and sucrose and was insensitive to 2-DG suggested that the Δ*col-26* mutant might have defects in sugar transport and/or metabolism. To test this hypothesis, we measured glucose uptake rates in WT, the Δ*cre*-1, and the Δ*col-26* mutants. Within the first 5 minutes, extracellular glucose was reduced to a similar level in all strains (Figure 7B), suggesting similar glucose transporting capacity. However, over the remaining 55 minutes, glucose uptake rates

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**Figure 6. Screen for function of new proteins involved in CCR.** (A) Growth assays of the WT, Δ*vib*-1, Δ*cre*-1, Δ*col-26*, Δ*creB*, and Δ*creD* strains on 2-deoxy-glucose (2-DG) when grown on 2% cellobiose VMM for 2 days or on 2% Avicel VMM for 4 days. (B) Effects of *col-26* and *cre-1* deletions on sensitivity to 2-DG and allyl alcohol. Strains were inoculated and grown in 2% cellobiose VMM with 100 mM allyl alcohol for 40 hrs. For 2-DG sensitivity tests, the strains were inoculated and grown in 2% Avicel with either 0.2% 2-DG or 0.5% 2-DG for 5 days.

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Figure 7. Deletion of col-26 causes defects in glucose sensing/metabolism. (A) Mycelial biomass of the Δcol-26 mutant relative to WT and the Δcre-1 strains on glucose, fructose, sucrose, or cellobiose. Mycelial biomass was measured at 24 hrs after inoculation. (B) Glucose uptake of WT, Δvib-1, Δcre-1, Δcol-26; Δvib-1 and the Δcre-1; Δcol-26; Δvib-1 mutants was assayed by monitoring glucose remaining in the medium at 5 min, 20 min, and 60 min from cultures of identical biomass.

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decreased dramatically in the Δcol-26 mutant (Figure 7B). These data indicate that the Δcol-26 mutant has defects in glucose sensing/metabolism, rather than in glucose transport.

Simultaneous inhibition of glucose sensing/metabolism and impairment of CRE1-mediated CCR rescues Δvib-1 growth on Avicel

Our data supported a role for CRE1 in CCR and a role for COL26 in the regulation of glucose utilization. We therefore tested sensitivity of the Δcre-1; Δvib-1 and the Δcol-26; Δvib-1 mutants to AA. The Δcre-1; Δvib-1 and the Δcre-1; Δcol-26 mutants were both sensitive to AA (Figure 8B), indicating the Δcre-1 mutation is epistatic for CCR to Δvib-1, while the Δcol-26; Δvib-1 mutant was insensitive to AA, consistent with the active CCR phenotype of the col-26 and the Δvib-1 mutants. However, although CCR was impaired in the Δcre-1; Δvib-1 mutant, the double mutant was still unable to produce cellulolytic enzymes and grow on Avicel (Figure 8A). Similar to the Δcol-26 mutant, the Δcol-26; Δvib-1 mutant also showed defects in glucose consumption (Figure 7B). Although the Δcol-26; Δvib-1 mutant was unable to utilize Avicel, it showed slightly higher enzyme levels than that of the Δvib-1 mutant (Figure 8A). We therefore hypothesized that simultaneously preventing CRE1-mediated CCR and reducing glucose sensing/metabolism via inactivation of col-26 would restore cellulase gene expression and enzyme activity in a vib-1 mutant.

As predicted, a Δcre-1; Δcol-26; Δvib-1 triple mutant utilized Avicel, produced significant cellulase activity and displayed a secretome similar to WT after 5 days of growth on Avicel (Figure 8A and S3). RT-PCR experiments from the Δcre-1; Δcol-26; Δvib-1 Avicel cultures showed that expression levels of clt-2 and cbb-1 were restored in the triple mutant (Figure 8B).

Although simultaneous deletion of cre-1 and col-26 restored utilization of cellulose in the Δvib-1 mutant, a significant lag in growth and enzyme activity in the triple mutant was observed as compared to the WT, Δcre-1, or Δcol-26 mutants (Figure 8C). To assess whether the Δcre-1; Δcol-26; Δvib-1 mutant was also delayed in transcriptional response upon exposure to cellulose, we measured expression levels of clt-2, cbb-1, cre-1, vib-1 and col-26 in the Δvib-1, Δcol-26, Δcre-1, Δvib-1; Δcol-26; Δvib-1, and Δcre-1; Δcol-26; Δvib-1 mutants as compared to the WT strain at 4 hrs and 24 hrs after cultures were shifted to Avicel conditions. Consistent with the enzyme activity assay and growth phenotype (Figure 8C), induction of clt-2 and cbb-1 was delayed in the Δcre-1; Δcol-26; Δvib-1 mutant (Figure 9). However, in the Δcol-26 mutant at the 4 hr time point, expression levels of cre-1 were significantly higher than in the Δvib-1 mutant, with the Δcol-26; Δvib-1 mutant showing an additive phenotype of significantly increased cre-1 expression levels. At the 24 hr time point, expression levels of cre-1 were only maintained in the Δvib-1 and Δcol-26; Δvib-1 mutants, but not in the Δcol-26 mutant. These data suggest that COL26 may function to repress cre-1 transcription to promote relief of CCR during the initial response to cellulolytic induction. Surprisingly, although the Δcre-1; Δvib-1 mutant was unable to utilize cellulose, induction of both clt-2 and cbb-1 were near WT levels at the 4 hr time point, unlike the Δvib-1 mutant (Figure 9A). However, at the 24 hr time point, expression levels of clt-2 were low and cbb-1 was undetectable in Δcre-1; Δvib-1 mutant (Figure 9B). These data suggest that although the Δcre-1; Δvib-1 can respond to cellulolytic induction by increasing clt-2 and thus cbb-1 expression levels, induction signaling cannot be maintained, perhaps due to repression by COL26 or by other factors present/absent in a Δvib-1 mutant background. The fact that the Δcre-1; Δcol-26; Δvib-1 mutant does not show WT restoration of initial cellulolytic induction (Figure 8C; Figure 9A) supports the hypothesis that additional unknown factors remain to be identified that play a role in nutrient sensing/signaling and the regulation of cellulose utilization in N. crassa.

Discussion

In this study, we showed that a Δvib-1 mutant displayed severe growth defects on cellulose, which was correlated with a lack of cellulolytic enzyme activity. By using RNA-seq data, we showed that expression of the Avicel regulon was significantly decreased in the Δvib-1 mutant, a phenotype that was rescued by constitutive expression of clt-2. Induction of clt-2 is dependent upon a signal cascade from cellulose or derivative and functional CLR1 (Figure 1) [8]. Here we showed that VIB1 is not involved in inducer signal processing or perception because the Δvib-1 mutant produced cellulolytic enzymes in response to cellulose. These data indicated that VIB1 functions upstream of regulators that mediate inducer-dependent signal transduction and cellulase gene expression and activity.

Our transcriptional profiling revealed that, under Avicel conditions, a deletion of vib-1 led to an increase in transcription of genes in metabolism and energy as well as genes reported to
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A. Cellulase activity

B. Relative mRNA expression

C. CMCase activity
mediate CCR. These results suggested that cellulolytic induction was mis-regulated in the \textit{D}vib-1 mutant. In the presence of glucose, \textit{N. crassa} adjusts its metabolism for a high rate of glycolysis and directs carbon flux to respiration and fermentation for biosynthesis and energy production [58], while genes involved in utilization of alternative carbon sources are repressed in a CRE1-dependent manner [26,27]. When lignocellulose is the only carbon source, CCR is relieved to allow the synthesis of “scouting” enzymes that liberate inducer molecules, such as cellobiose [9,44,59]. In \textit{S. cerevisiae}, glucose is sensed through a multifaceted mechanism including direct detection of glucose by glucose receptors/transporters on the plasma membrane and by the sensing of glucose-6-P and other metabolites by metabolic enzymes. The glucose signals are transmitted to CCR mainly through the Snf1 complex and the Mig1 (CreA/Cre1 homolog) transcriptional repressor complex [60,61]. In \textit{A. nidulans}, mutations in two hexose kinase genes (\textit{hxkA/glkA4}) results in inappropriate de-repression of genes under glucose growth.

**Figure 8. Simultaneous deletion of \textit{cre-1} and \textit{col-26} rescues the phenotype of \textit{D}vib-1 on cellulose.** (A) Cellulase activity of culture supernatants after 4-days of growth on Avicel from WT versus \textit{D}vib-1, \textit{D}cre-1, \textit{D}col-26, \textit{D}cre-1; \textit{D}vib-1, \textit{D}col-26; \textit{D}cre-1; \textit{D}vib-1 strains. (B) RT-PCR measurements of \textit{clr-2} and \textit{cbh-1} expression in the WT versus the \textit{D}col-26, \textit{D}cre-1, and the \textit{D}cre-1; \textit{D}col-26; \textit{D}vib-1 cultures after 5-days of growth on Avicel. Expression levels were normalized to WT. (C) The CMCase activity of Avicel cultures of WT versus \textit{D}col-26, \textit{D}cre-1, and the \textit{D}cre-1; \textit{D}col-26; \textit{D}vib-1 mutants during a time course of growth on Avicel.

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**Figure 9. Suppression of \textit{cre-1} and \textit{col-26} expression by VIB1 plays a role in early inductive and utilization phase during growth on cellulose.** The transcriptional expression of \textit{cre-1}, \textit{vib-1}, \textit{col-26}, \textit{clr-2}, and \textit{cbh-1} were measured by RT-PCR at 4 hrs (A) and 24 hrs (B) after 16 hr sucrose growth cultures were transferred to Avicel conditions. Expression levels were normalized to WT.

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conditions, although to a lesser extent than a creA mutant strain [29]. Here we show that simply eliminating CRE1-mediated CCR did not rescue the growth defect of Δvib-1 mutant on Avicel, but that a deletion of col-26 was also required.

The Acol-26 mutant exhibited a growth defect on glucose, fructose and sucrose, which was not associated with a deficiency in glucose transport (Figure 7B). In T. reesei, a strain carrying a mutation in bglR shows reduced expression of β-glucosidase genes, suggesting the BglR plays a positive role in CCR by increasing glucose release from cellulose [15]. However, our analyses of cellulolytic activity of secreted enzymes in the Acol-26 mutant showed no difference in glucose versus cellobiose release (Figure 8A), a result that is in contrast to the strongly reduced glucose release from culture supernatants in the Δ3βG mutant (which lacks extracellular β-glucosidase activity) (Figure S2). Although we have not determined how glucose metabolism is changed in the Acol-26 mutant, the resistance of Acol-26 to 2-DG inhibition suggests a defect in glucose sensing/metabolism; CRE1-mediated CCR was still functional (as shown by insensitivity to AA). The fact that a deletion of col-26 and cre-1 restored growth of Δvib-1 on Avicel suggests a synergistic effect between glucose sensing/metabolism mediated by COL26 and CRE1-regulated CCR in repressing cellulolytic induction (Figure 10). However, other unknown factors in addition to CRE1 and COL26 play a role in the Δvib-1 mutant. The Δcre-1; Acol-26; Δvib-1 mutant showed a significant lag in gene induction and enzyme secretion under cellulolytic conditions (Figure 8C). Future experiments to identify additional mutations that fully suppress the Δvib-1 cellulolytic phenotype and the identification of direct targets of VIB1 will be most informative for further dissection of glucose sensing and CCR in filamentous fungi.

Our data supports the model that the regulatory function of VIB1 on CRE1-mediated CCR and COL26-mediated glucose sensing/metabolism functions during different stages of the cellulolytic response (Figure 10). At induction stage, both VIB1 and COL26 negatively regulate CRE1-mediated CCR (Figure 9), thus allowing a relief of CCR and efficient induction of cellulolytic genes in response to cellulose. During the utilization phase, glucose is released from cellulose, and glucose sensing/signaling via COL26 may repress cellulolytic responses, with VIB1 functioning to dampen this inhibition. As many cellulolytic genes are subject to carbon catabolite repression and a requirement for CLR2 for induction, the cellular response to plant biomass may depend on the relative strength of these two antagonizing forces (Figure 10). Mechanistically, how VIB1 exerts its function on glucose sensing/metabolism via COL26 and CCR via CRE1 remain to be elucidated.

In the hyper-secreting T. reesei strain, RUT-C30, disruption of phosphoglucose isomerase gene (pgi1) blocks formation of fructose-6-P from glucose-6-P and increased cellulase production on glucose. This increase relies on a genetic interaction between the Δpgi1 mutation and the cre1-1-1 mutation in the RUT-C30 background [62]. Interestingly, both the hyper-secreting T. reesei RUT-C30 and PC-3-7 strains have mutations in cre1 and bglR/col-26 [15,63,64], but whether a synergy exists between Δcre1 and ΔbglR in T. reesei, as in N. crassa, and its relationship to T. reesei vib1 is unclear. Many cellulolytic enzyme hyper-producers such as T. reesei RUT-C30 and PC-3-7, and P. decumbens JU-A10-T show relief from CCR, but contain a large number of mutations in additional genes that contribute to the hyper-production phenotype [2,15,63–65]. Identifying and characterizing possible synergistic effects of the different mutations on hyper-production of lignocellulose enzymes, as shown in this study, will be a challenge.

The function of VIB1 in regulating glucose sensing/metabolism and CCR plays a role in the utilization of other complex substrates. VIB1 is required for extracellular protease production in response to carbon and nitrogen starvation, a function shared by its homolog in A. nidulans, xprG [66–70]. The Δvib-1 mutant also exhibits inappropriate temporal and spatial conidiation and has defects in protoperithecia formation [48,70], two developmental events that are regulated by nitrogen and glucose limitation and signaling [71]. A shotgun proteomic analysis of culture

Figure 10. Model for role of VIB1 in regulating glucose sensing/metabolism and CCR under cellulolytic conditions. In an early encounter of N. crassa to cellulose, “scouting” enzymes induced by carbon starvation act on cellulose to liberate cellulolytic inducers that activate signaling cascades that include activation of CLR1, and subsequent expression of CLR2 and induction of genes encoding cellulases. Efficient cellulolytic induction also requires de-repression of carbon catabolite repression (CCR), as many cellulase genes are not expressed even in the presence of an inducer if a preferred carbon source is available. In the early cellulolytic induction, VIB1 functions to repress CRE1-mediated CCR and glucose sensing/metabolism by COL26, which results in CCR de-repression and productive cellulolytic responses. COL26 also plays a role in sensing cre-1 expression and thus alleviating CCR. As cellulases are produced, glucose is liberated from cellulose and N. crassa transits into the utilization phase, which is associated with reduced transcription of cellulases [44]. Repressive function of VIB1 on col-26 expression may be important for tuning cellular responses for the need to produce sufficient enzymes to liberate simple sugars from cellulose, but without over-activating CCR. VIB1 also likely regulates other factors important for cellulolytic induction.

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supernatant of the Pshb-1 strains under carbon source depletion showed a higher amount of intracellular proteins relative to WT (Table S3). These data are in consistent with a role of VIB1 in promoting cell death [47,48,72], and in autolysis in A. nidulans [73], perhaps via perturbed nutritional signaling. Autolysis is frequently observed in submerged batch cultures in industrial bioprocessing, and promotes cryptic growth for survival and protein production under nutrient-depleted conditions [74]. Further manipulations of vib-1 and its homologs in filamentous fungi may yield economic benefits via the regulation of autolysis under industrial settings.

In summary, our data show that VIB1 is an essential regulator for cellulase production under inductive conditions and identifies COL26 as an important player in glucose sensing/metabolism. As VIB1 mediates metabolic changes as well as programmed death, two properties shared by mammalian tumor suppressor p53 [75,76], the molecular mechanism in linking the two could be conserved, and further investigation of vib-1 function and its homologs in filamentous fungi may also shed light on cancer research.

Materials and Methods

Strains

FGSC 2489 was used as the WT reference strain and background for mutant strains [46]. FGSC 11308 (Δvb-1; mat a), FGSC 11309 (Δvb-1; mat A), FGSC 11030 (Δcol-26; mat a), FGSC 11031 (Δcol-26; mat A) were obtained from the Fungal Genetics Stock Center (http://www.fgsc.net/) [50]. The vib-1 mis-expression strain Pshb-1 (Pshb-1; Δvb-1) was constructed by transforming FGSC 11308 with a DNA fragment containing the promoter of the clock controlled gene 1 (ccg-1) and the open reading frame and 3' untranslated region (UTR) of vib-1 and homologous and flanking regions from the coding sequence of the his-3 gene. Transformants were selected for histidine prototrophy [77] and backcrossed to FGSC 2489 to obtain a his-3::ccg-1·vib-1; Δvb-1·homokaryotic strain. The Ty vib1 mis-expression strain PTv-vib1 (PTv vib1; Δvb-1) was created in the same way except that the open reading frame and 3'UTR of Tr vib1 was used. The Pc cl-2, the Acre-1, the Δβ3G and the Δβ3G Acre-1 strains were from previous studies [9,10,26]. The P cl-2; Δvb-1 strain, the Δβ3G; Δvib-1 strain, the Δcol-26; Δvib-1 strain, the Acre-1; Δcol-26 strain, and the Acre-1; Δcol-26; Δvb-1 strain were created through crosses.

Culture conditions

N. crassa cultures were grown on Vogel's minimal medium (VMM) [78]. Unless noted, 2% (w/v) sucrose was used as a carbon source. Strains were pre-grown on 3 mL VMM slants at 30°C in dark for 24 hrs, then at 25°C in constant light for 4–10 days to stimulate conidia production. For flask cultures, conidia were inoculated into 100 mL of liquid media at 10⁶ conidia/mL and grown at 25°C in constant light and shaking (200 rpm). To test 2-DG and allyl alcohol sensitivity, 3 mL of liquid media containing either 0.2% (w/v) 2-DG (Sigma Aldrich, MO) or 100 mM allyl alcohol were inoculated with 10⁶ conidia/mL and grown in 24-well plates at 25°C in constant light and shaking (200 rpm).

For crosses, one parental strain was grown on synthetic crossing medium [79] as the female for 2 weeks at room temperature for protoperithecia development. The other parental strain was used as the male to fertilize the protoperithecia. Crosses were kept for 3 weeks at room temperature. Ascospores were collected and activated as described [80], plated on 1% VMM, and incubated at room temperature for 18 hrs. Germinated ascospores were selected and transferred to selective slants for further screen and confirmation.

Media shift experiments

Cultures were grown on sucrose for 16 hrs, centrifuged at 2000 g for 10 min and washed in VMM or MM without a carbon source, followed by 4 hrs growth in 100 mL VMM or MM with 2% carbon source (sucrose, cellobiose, Avicel PH-101 (Sigma Aldrich, MO)) or with no carbon source added.

RNA preparation and qRT-PCR analysis

Mycelia were harvested by filtration and flash frozen in liquid nitrogen. RNA was extracted using the Trizol method (Invitrogen) and further purified using RNeasy kits (QIAGEN). Four ng of RNA was used as template in each quantitative RT-PCR (qRT-PCR) reaction. qRT-PCR was carried out using EXPRESS One-Step SYBR GreenER kit (Invitrogen) and Applied Biosystems Step One Plus Real Time PCR system. qRT-PCR were done in biological duplicates or triplicates with actin as the endogenous control. Relative expression levels were normalized to actin, and fold changes in RNA level were the ratios of the relative expression level on inducing conditions to no carbon conditions.

RNA sequencing and transcription expression analysis

Libraries were prepared according to standard protocols from Illumina Inc (San Diego, CA) and sequenced on the HiSeq 2000 platforms at QB3 Vincent J. Coates Genomics Sequencing Laboratory (CA). Sequenced reads were mapped against predicted transcripts from the N. crassa OR7A4 genome [81]/Neurospora crassa Sequencing Project, Broad Institute of Harvard and MIT http://www.broadinstitute.org/) with Tophat v2.0.4 [82]. Transcript abundance (FPKM) was estimated with Cufflinks v2.0.2 mapping against reference isoforms and differential gene expression were analyzed with Cuffdiff v2.0.2 [83]. Biological replicates used for RNA-seq showed high reproducibility. The Pearson correlation of FPKM on log basis (p-value<2.2e-16): rp ≥ 0.96 between WT (Nc) replicates, rp ≥ 0.91 between WT (Av) replicates, rp ≥ 0.99 between Δvb-1 (Nc) replicates, and rp ≥ 0.96 between Δvb-1 (Av) replicates.

For hierarchical clustering analysis, FPKM were log transformed, normalized and centered on a per gene basis with Cluster 3.0 [84] so that values from each gene ranged from −1 (minimum) to 1 (maximum). Average linkage clustering was performed with Euclidean distance as the similarity metric. Functional category analysis was done as described in [8]. Lists of genes were matched against the MIPS Functional Category Database [85], and significance of enrichment was calculated.

Enzyme activity assays

For CMCase and xylanase activity assays, Azo-CM-Cellulose and Azo-xylan (Beechwood) from Megazyme (Wicklow, Ireland) were used as substrates. Protein concentration was measured with the Bradford assay (BioRad). Cellulase assays were conducted by mixing 500 µL of culture supernatant with 500 µL 0.5% (w/v) Avicel in 100 mM sodium acetate, pH 5.0, and incubated with shaking at 37°C for 5 hrs. Reactions were stopped by centrifugation at 2000 g for 5 min and by addition of 9 volumes of 0.1 M NaOH to the reaction supernatants. Released glucose and cellobiose were separated on a PA-200 HPAEC column and analyzed on Dionex ICS-3000 as described in [45].

Glucose uptake assays

For CMCase and xylanase assays, Azo-CM-Cellulose and Azo-xylan (Beechwood) from Megazyme (Wicklow, Ireland) were used as substrates. Protein concentration was measured with the Bradford assay (BioRad). Cellulase assays were conducted by mixing 500 µL of culture supernatant with 500 µL 0.5% (w/v) Avicel in 100 mM sodium acetate, pH 5.0, and incubated with shaking at 37°C for 5 hrs. Reactions were stopped by centrifugation at 2000 g for 5 min and by addition of 9 volumes of 0.1 M NaOH to the reaction supernatants. Released glucose and cellobiose were separated on a PA-200 HPAEC column and analyzed on Dionex ICS-3000 as described in [45].
cultures were thoroughly washed with MES buffer (10 mM 2-(N-morpholino)ethanesulfonic acid, 100 mM NaCl), and each washed culture was transferred into 4 ml of MES buffer supplemented with 10 mM glucose and grown at room temperature for 1 hr with shaking at 350 rpm. Culture supernatants were sampled at 5, 20, and 48 hr, and diluted in 53 volumes of 0.1 M NaOH. Glucose levels were measured using Dionex ICS-3000 HPAEC-PDA 200 and MES buffer instead of VMM was used to avoid precipitation that interferes with downstream analysis.

Protein gel electrophoresis
Culture supernatants were mixed with 4× SDS loading buffer and boiled for 10 min before loading onto Criterion 4–15% Tris-HCl Precast Gel (Bio-Rad). GelCode Blue Stain Reagent (Thermo Scientific) was used for gel staining.

Microscopy and imaging
Strains were inoculated in 2% sucrose VMM and grown at 25°C for 12 hr in eight-chamber Lab-Tek chambered cover glasses (Nalge Nunc International, Naperville, IL). Localization of VIB1-GFP was observed using a 100×1.4 NA oil immersion objective on a Leica SD6000 spinning disk confocal with 488 nm laser and controlled by Metamorph software. Z-series stacks were collected and maximum intensity projections were created using ImageJ. For medium shift experiment, the cultures in the chamber were washed with VMM without carbon sources and VMM with 0.5% Avicel was added, followed by immediate time-lapse recordings with an interval of 15 min.

Proteomic analysis
Equal volume of culture supernatants of WT and P. stipitis vib-1 mutants was subjected to SDS-PAGE and secretome proteins identified as described in [86]. In-gel trypsin-digestion was performed according to manufacture protocol (Promega, Trypsin Gold). Digested peptides were separated using ProtID-Chip-43 (II) and analyzed using the Agilent 6510 Q-TOF LC/MS as in [9].

Supporting Information

Figure S1 VIB1 functions in xylanase production and localizes to nuclei in both sucrose and cellulose media. (A) The vib-1 mutant grew slowly on xylan and showed reduced xylanase activity. (B) Mycelial biomass accumulation after 24 hrs of growth in the Δvib-1 mutant as compared to WT in VMM containing 2% (w/v) of glucose, cellulose, or xylene as the sole carbon source. (C) Fluorescence microscopy showing localization of VIB1-GFP to nuclei under both sucrose and Avicel conditions.

Figure S2 Mutations in ΔβG genes rescue the cellulase-deficient phenotype of the Δvib-1 mutant on Avicel. Cellulase activity in the ΔβG; Δvib-1 mutant compared to the WT, Δvib-1 and the ΔβG strains after 4 days of growth on 2% Avicel. Cellulase activity was measured using Avicel as substrate (see Materials and Methods).

Table S1 Gene list of the 91 Avicel-regulon genes differentially expressed in Δvib-1 as compared to WT on Avicel.

Table S2 Gene list of the 770 genes differentially expressed between no carbon and Avicel conditions in Δvib-1.

Table S3 Intracellular N. crassa proteins enriched in the supernatant of vib-1 cultures relative to WT after carbon depletion.

Author Contributions
Conceived and designed the experiments: YX JS NLG. Performed the experiments: YX JS. Analyzed the data: YX JS NLG. Contributed reagents/materials/analysis tools: NLG. Wrote the paper: YX NLG.

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