

Biological nitrification inhibition (BNI) in *Brachiaria* pastures: A novel strategy to improve eco-efficiency of crop-livestock systems and to mitigate climate change

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Keywords: Ammonium-oxidizing microorganisms, nitrous oxide emissions, nitrogen use efficiency, genetic variation.

Introduction

Up to 70% of the nitrogen (N) fertilizers applied to agricultural systems are lost due to nitrification and denitrification. Nitrification is a microbiological process that generates nitrate (NO₃⁻) and promotes the losses of N fertilizers by leaching and denitrification. Nitrification and denitrification are the only known biological processes that generate nitrous oxide (N₂O), a powerful greenhouse gas contributing to global warming. There is an urgent need to suppress nitrification process in soil to improve N-recovery and N use efficiency (NUE) of agricultural systems and to mitigate climate change (Subbarao *et al.* 2012). Certain *Brachiaria* grasses (*B. humidicola*) can suppress soil-nitrification by releasing biological nitrification inhibitors (BNIs) from roots, thereby reducing N₂O emissions. This phenomenon, termed biological nitrification inhibition (BNI), has been the subject of recent research to characterize and validate the concept under field conditions (Subbarao *et al.* 2009). Advances on three aspects of BNI research are reported here: (1) gene quantification of soil nitrifying microorganisms to determine BNI activity in *B. humidicola*; (2) screening of *B. humidicola* breeding materials to identify hybrids with contrasting levels of BNI; and (3) quantification of the BNI-residual effect from *B. humidicola* on N-recovery and agronomic-NUE of the subsequent maize crop.

Methods

Gene quantification of soil nitrifying microorganisms to determine BNI activity in B. humidicola

A proof of concept work was designed to monitor the dynamics of nitrification in soils as influenced by *Brachiaria* spp. with differential BNI capacities (Subbarao *et al.* 2009). Soybean crop and bare soil, which lack such BNI capacity, were used as controls. Ammonium-sulfate was applied to each plot. Copy number of *amoA* genes of ammonia-oxidizing bacteria (AOB) and archaea (AOA) were determined through Real-Time PCR to quantify the

impact of inhibitory effect from *Brachiaria* sp. under field conditions at 1 day after the ammonium-sulfate application.

Screening of B. humidicola breeding materials to identify hybrids with contrasting levels of BNI

A set of apomictic *B. humidicola* hybrids were screened by determining nitrification rates in soil samples taken from unreplicated field plots established for seed production. Four CIAT accessions were used as controls for BNI activity.

Quantification of the BNI-residual effect from B. humidicola on N-recovery and NUE of the subsequent maize crop

One-hectare of field area was selected from each of the three contrasting land uses that included a long-term *B. humidicola* CIAT 679 pasture (15-year-old) with accumulated inhibitory effect in soil (*i.e.*, high BNIs in soil), a nearby agricultural field (in which a crop rotation of maize and soybean was practiced for 4 years) with low BNIs in soil, and a native savanna field with moderate level of BNIs in soil. These three sites were considered as land use treatment sites. Maize hybrid (Pioneer 30K73) was sown on 17 July 2012 in all three field sites. Nitrogen fertilizer was applied at three rates (60, 120 and 240 kg/ha) at each field site. Grain yield and agronomic NUE were determined to assess the BNI residual effect on subsequent maize cultivation.

Results

Molecular data confirmed that *B. humidicola* CIAT 16888 has the capacity to inhibit soil nitrification (BNI activity). Rhizosphere soil from *B. humidicola* CIAT 16888 plots exhibited a lower gene copy number of AOB and AOA *amoA* genes compared to the controls (soybean and bare soil) and the other tropical grasses (Fig. 1). Different values of nitrification rates observed in field plots of *B. humidicola* breeding materials suggested genetic variation for BNI and contributed to identification of hybrids with

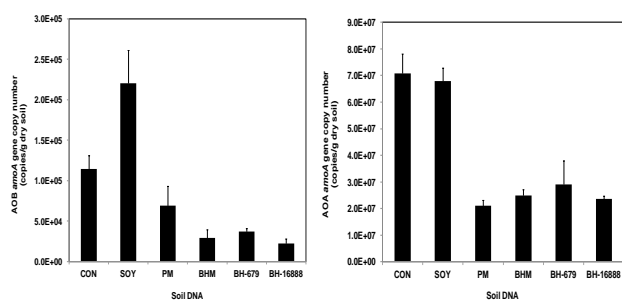


Figure 1. Gene copy number of ammonia-oxidizing bacteria (AOB) *amoA* gene (left), and ammonia-oxidizing archaea (AOA) *amoA* gene (right) at 1 day after ammonium-sulfate application. CON = control (bare soil); SOY = soybean; PM = *Panicum maximum*; BHM = *Brachiaria* hybrid cv. Mulato; BH-679 = *B. humidicola* CIAT 679 (standard cultivar); BH-16888 = *B. humidicola* CIAT 16888 (a high-BNI capacity germplasm accession). Gene copy number was expressed as copy number per g of dry soil. Values are mean ± SE from three replications

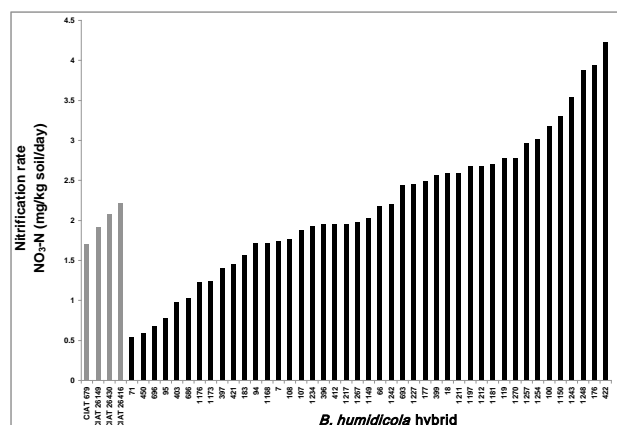


Figure 2. Genotypic differences in nitrification rates – expressed as NO₃-N (mg/kg soil/day) in field plots of *B. humidicola* hybrids. Gray bars represent *B. humidicola* CIAT accessions used as controls

contrasting BNI capacities (Fig 2). The higher grain yields of maize observed from *B. humidicola* pasture land use were associated with greater values of agronomic NUE, particularly at lower rate of N applied (60 kg/ha). This observation indicates the importance of accumulated BNIs from this pasture over time in improving the agronomic-NUE of maize crop (Fig. 3).

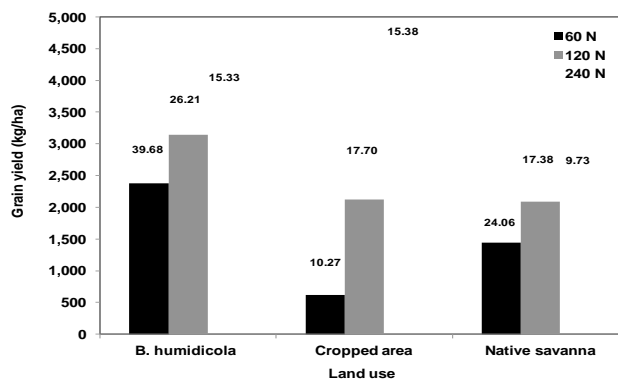


Figure 3. Grain yield (kg/ha) from maize plots fertilized with 60, 120, and 240 kg N/ha where a long term (15 years-old) *B. humidicola* pasture, cropped area (maize rotated with soybean), and native savanna were previous land use systems. Agronomic nitrogen use efficiency (kg of grain yield/kg of N applied) values are shown above the SE bars. Values are means ± SE from three replications

Conclusion

BNI activity in *Brachiaria humidicola* plots was confirmed by observing a lower copy number of *amoA* genes from bacterial and archaeal populations compared to soybean and bare soil plots. The wide variation of in nitrification rates observed in a set of apomictic *B. humidicola* hybrids contributed to identification of hybrids with contrasting BNI capacities. Accumulation of BNIs in soil of long-term *B. humidicola* pasture improved grain yield and agronomic-NUE of the subsequent maize crop.

References

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