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### Insufficient nitrogen supply from symbiotic fixation reduces seasonal crop growth and nitrogen mobilization to seed in highly productive soybean crops

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#### **ORIGINAL ARTICLE**

### WILEY

### Insufficient nitrogen supply from symbiotic fixation reduces seasonal crop growth and nitrogen mobilization to seed in highly productive soybean crops

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#### Abstract

Nitrogen (N) supply can limit the yields of soybean [Glycine max (L.) Merr.] in highly productive environments. To explore the physiological mechanisms underlying this limitation, seasonal changes in N dynamics, aboveground dry matter (ADM) accumulation, leaf area index (LAI) and fraction of absorbed radiation (fAPAR) were compared in crops relying only on biological N<sub>2</sub> fixation and available soil N (zero-N treatment) versus crops receiving N fertilizer (full-N treatment). Experiments were conducted in seven high-yield environments without water limitation, where crops received optimal management. In the zero-N treatment, biological N2 fixation was not sufficient to meet the N demand of the growing crop from early in the season up to beginning of seed filling. As a result, crop LAI, growth, N accumulation, radiation-use efficiency and fAPAR were consistently higher in the full-N than in the zero-N treatment, leading to improved seed set and yield. Similarly, plants in the full-N treatment had heavier seeds with higher N concentration because of greater N mobilization from vegetative organs to seeds. Future yield gains in high-yield soybean production systems will require an increase in biological N<sub>2</sub> fixation, greater supply of N from soil or fertilizer, or alleviation of the trade-off between these two sources of N in order to meet the plant demand.

#### KEYWORDS

Glycine max (L.) Merr., leaf area, nitrogen, soybean, symbiotic fixation

### 1 | INTRODUCTION

Soybean [*Glycine max* (L.) Merr.] is the world's most important legume crop and the primary vegetable protein source for food and feed (Wilson, 2008). Its nitrogen (N) requirement per unit of photosynthate

produced is the highest among all food crops (Sinclair & de Wit, 1975). On average, soybean requires 80 kg N in aboveground dry matter (ADM) per metric ton of seed produced (Salvagiotti *et al.*, 2008; Tamagno *et al.*, 2017). This value is about three times greater than the N requirement per unit of grain in cereal crops such as rice, wheat, or

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maize (Barraclough *et al.*, 2010; Setiyono, Walters, Cassman, Witt, & Dobermann, 2010a; Yin *et al.*, 2019). Unlike cereal crops, fertilizer N is rarely used on soybean, except for an occasional small amount applied as 'starter' at sowing. Hence, soybean relies on two major N sources to meet its large N requirement: (a) biological  $N_2$  fixation (BNF) and (b) N from soil organic matter mineralization, inorganic soil N left by previous crop, atmospheric dry and wet deposition, water table and irrigation water (hereafter collectively called 'indigenous soil N supply'). However, it is unclear if N from BNF and indigenous soil N supply are sufficient to meet soybean N requirement as average yield continues to increase due to improved cultivars and agronomic practices (Grassini, Specht, Tollenaar, Ciampitti, & Cassman, *et al.*, 2014; Specht *et al.*, 2014).

There is a well-known trade-off between BNF and indigenous soil N supply, that is, BNF decreases as the contribution from indigenous soil N supply increases and vice versa (Santachiara, Borrás, Salvagiotti, Gerde, & Rotundo, 2017; Streeter & Wong, 1988). As a result, application of fertilizer N reduces BNF, resulting in a small marginal increase in crop N uptake and vield (Salvagiotti et al., 2008). While this trade-off has often been reported, it is not clear whether the combined N supply from BNF and soil is sufficient to meet the N requirement in soybean in highly productive environments (Ciampitti & Salvagiotti, 2018). To address this question. Cafaro La Menza, Monzon, Specht, and Grassini (2017); Cafaro La Menza et al. (2019) developed a protocol to assess N limitation across a wide range of environments, where seed yields ranged from 2.5 to 6.7 Mg ha<sup>-1</sup>. The protocol consisted of a side-byside comparison between a 'control' treatment (hereafter called 'zero-N') that forced the crop to rely on BNF and indigenous soil N supply, and a 'full-N' treatment designed to provide the crop with fertilizer N supply to optimally match the expected seasonal plant N demand. The full-N treatment aimed to eliminate N limitation at any time of the growing season, independent of the contribution from BNF. Average seed yield was higher in the full-N than in the zero-N treatment, but the magnitude of the yield difference was larger in highly productive environments, confirming the existence of a N limitation in high-yield soybean.

The studies by Cafaro La Menza et al. (2017, 2019) also documented that higher seed yields in the full-N versus zero-N treatment were associated with greater end-of-season shoot dry matter, total N, seed number and seed mass. Despite the other well-known trade-off between seed yield and seed N concentration (Chung et al., 2003), it was remarkable that the full-N treatment also exhibited higher seed N concentration, which is of interest for soybean processors who seek high protein concentration when producing meal for animal feed (Brumm & Hurburgh Jr, 2006). In contrast, there were no differences in harvest index, N-use efficiency, N-harvest index or seed oil concentration between the full-N and zero-N treatments. While these prior studies advanced knowledge of N limitation in high-yield soybean crops, they did not provide insight into the physiological mechanisms over a seasonal timeframe that eventually led to the differences in seed yield and seed N concentration. Such an assessment would be useful for a more mechanistic understanding of the N limitation in high-yield soybean production systems.

Considering that soybean yield has to increase to meet future demand while avoiding further conversion of natural ecosystems into

cropland (Cassman & Grassini, 2020), and that future yield gains should not have negative effects on seed protein concentration, it is important to design strategies to overcome the N limitation in highyield soybean systems. With that goal in mind, the objective of this study was to better understand seasonal N dynamics by which key physiological mechanisms (*e.g.*, BNF, ADM and N accumulation, leaf area index (LAI), photosynthesis, and N mobilization) account, separately or in concert, for the observed differences in seed yield and seed N concentration between soybean crops growing under contrasting N supply. For that comprehensive assessment of N limitation in soybean, we used original data on accumulated ADM and N, BNF and other physiological processes collected during the entire crop season from field experiments conducted over 2 years in a high-yield environment.

#### 2 | MATERIALS AND METHODS

#### 2.1 | Experimental sites and design

This article presents new data of seasonal dynamics of ADM, N, LAI, and other physiological variables from the set of experiments conducted in Nebraska (NE), USA by Cafaro La Menza *et al.* (2017, 2019) (Table 1). These previous studies addressed the occurrence of N limitation in soybean at different seed yield levels and the influence of indigenous soil N supply on the magnitude of N limitation by looking at end-of season productivity and total N. Here, we investigated the underlying mechanisms that explain the nature of the N limitation in high-yield production systems by looking at original data collected during the growing season from the same set of experiments. These NE irrigated production systems provide a suitable background to evaluate N limitation in high-yield soybean, considering that NE has the largest irrigated soybean area in the world (*ca.* 2.6 million ha) where producers achieve average irrigated yields of *ca.* 4.4 Mg ha<sup>-1</sup> (USDA-NASS, 2014–2018; https://www.nass.usda.gov/).

Experiments were conducted in pivot-irrigated producer fields located at four sites across NE (Mead, Saronville, Smithfield and Atkinson) during 2016 and 2017. All fields at these sites have consistently produced high soybean yields (>5 Mg  $ha^{-1}$ ) in previous years. This combination of site-years portrayed well the range of weather, soil, and management (e.g., tillage methods, cultivar maturity groups [MGs]) used in NE and the rest of the US North-Central region (Table 1). We excluded the experiment in Atkinson in 2016 due to a severe infestation of powdery mildew (Microsphaera diffusa). A portable weather station was installed within 50 m of each experiment in both years to monitor hourly solar radiation, soil and air temperature, rainfall, relative humidity and wind speed. A well-validated soybean crop model (SoySim; Setiyono et al., 2010b) was used to simulate the yield potential for each site-year using local weather and recorded sowing date and cultivar MG at each site-year (Table 1). In all cases, simulated yields exceeded the yield threshold of 4.5 Mg ha<sup>-1</sup> proposed in the literature to define high-yield soybean production environments where N limitation is likely to occur (Salvagiotti et al., 2008).

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Description of the seven field experiments conducted in Nebraska, USA

**TABLE 1** 

Site	Crop season	Cultivar (and MG)	Sowing date	Seeding rate (m <sup>-2</sup> )	Yield potential (Mg ha <sup>-1</sup> )	Tillage method	pHa	soc <sup>a</sup> (g kg <sup>-1</sup> )	Soil P <sup>a</sup> (mg kg <sup>-1</sup> )	Soil K <sup>a</sup> (mg kg <sup>-1</sup> )	Applied fertilizer (kg ha <sup>-1</sup> ) <sup>b</sup>
Mead (41.244 N; 96.502 W)	2016	AG2723 (MG 2.7)	May 8	25	6.4	Strip till	6.3	22	12	335	N (20); P (21); S (17); Zn (2)
	2017	AG2723 (MG 2.7)	May 6	35	6.4	Strip till	6.7	24	25	268	N (12); P (25)
Saronville (40.601 N; 97.966 W)	2016	AG2431 (MG 2.4)	April 26	25	5.5	Disk	6.2	21	17	395	N (17); P (35); S (10); Zn (1)
	2017	P33T72R (MG 3.3)	April 23	35	5.6	Strip till	7.1	20	33	438	N (6); P (20)
Smithfield (40.538 N; 99.683 W)	2016	P24T19 (MG 2.4)	May 13	32	5.7	No till	7.5	16	12	487	P (38); S (1)
	2017	P27T59 (MG 2.7)	May 25	44	4.8	No till	7.2	18	25	373	N (24); P (48); S (27)
Atkinson (42.658 N; 99.029 W)	2017	AG2723 (MG 2.7)	April 24	40	6.3	Disk	7.0	14	7	184	N (20); P (22); K (50); Ca (62); Mg (8); S (33); Zn (2
Vote: K: 1 M NH <sub>4</sub> -acetate extrac	stable potas	sium; P: Bray-1 pho	sphorus; N	lG: maturity gr	oup; SOC: soil orga	nic carbon; S: s	ulfur; (	a: calciun	n; Mg: magnes	sium; Zn: zinc.	

Average concentration in the 0.60 m depth measured at crop emergence (VE stage)

Parenthetic values indicate rates of elemental nutrient applied to both treatments (full-N and zero-N) at or before sowing in each environment.

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Each of the seven site-year combinations is referred to hereafter as an 'environment' that included two N treatments (full-N and zero-N) in a complete randomized design with four replicates per treatment (size: 176 m<sup>2</sup> each). Experimental plots were purposely placed in areas within each field where the highest (maize and/or soybean) yields had been achieved in previous years. Experiments were conducted in fields following a 2-year rotation with maize, which is the dominant practice in the US North Central region (Grassini et al., 2014). The practice of coating seeds with rhizobia inoculum (commonly referred to as 'inoculation') is not needed for proper nodulation in fields with prior soybean history as documented by a number of field studies (e.g., Carciochi et al., 2019; de Bruin et al., 2010; Leggett et al., 2017 and references cited therein). Hence, in our experiments, seeds were treated with fungicide and insecticide, but received no inoculant. Row spacing was 0.76 m in all environments, with seeding rates adjusted to maximize soybean yields (De Bruin & Pedersen, 2009). Soil water content in the upper meter was monitored using Watermark® sensors (Irmak, Payero, VanDeWalle, Rees, & Zoubek, 2014), and maintained above 50% of plant available water throughout the entire growing season, except for a short period of time at Atkinson in 2017 (Supplementary Figure S1). Several prophylactic foliar applications of herbicide, fungicide and insecticide and presowing nutrient applications (based on soil test results) kept the crops free from biotic and nutrient stresses (aside from N in the zero-N treatment). Inorganic soil N in the upper 0.60 m at sowing ranged from 25 to 58 kg N ha<sup>-1</sup> across environments, which is within the range expected for soybean grown after a maize crop (Farmaha et al., 2016).

#### 2.2 Nitrogen treatments

Two N treatments were compared in each environment: (a) a 'zero-N' treatment in which the crop relied on indigenous soil N supply and BNF and (b) a 'full-N' treatment designed to provide the crop with sufficient fertilizer N to optimally match seasonal crop N demand. The N fertilizer was applied as urea and broadcast between plant rows in the full-N treatment. A total seasonal amount of 870 kg N ha<sup>-1</sup> was applied at all sites based on (a) site-specific yield potential simulated using the SoySim model (Setiyono, et al., 2010b), (b) N uptake requirement of 80 kg N per Mg<sup>-1</sup> seed yield (Salvagiotti et al., 2008; Tamagno et al., 2017), and (c) an extra 40% of fertilizer N to compensate for potential N losses through volatilization and leaching resulting from mismatches between irrigation or rainfall events and fertilizer N application. Total N fertilizer amount was split into five applications to approximate the expected increase in crop N requirement during the crop season (Bender, Haegele, & Below, 2015; Thies, Singleton, & Bohlool, 1995). Of the total N fertilizer amount, 10%, 10%, 20%, 30%, and 30% were applied at the V2, V4, R1, R3 and R5 stages, respectively (Figure 1).

#### 2.3 **Field measurements**

Detailed measurements of phenology, leaf area, photosynthesis and accumulated ADM and N were made at regular intervals over the entire





crop season. Phenological stages defined by Fehr & Caviness (1977) (Figure 1) were recorded from 10 consecutive plants within one row in each replicate for both treatments every seven days, starting at VE and concluding at R7. Following Lindquist *et al.* (2005), phenological stages were made comparable across environments (with different temperature regimes, sowing date and cultivar MG) by defining each development stage (DS) using a dimensionless scale where 0, 1 and 2 correspond to the VE, R3 and R7 stages, respectively (Figure 1). In our case, each DS was calculated based on daily mean air temperature using a beta function as defined by Wang & Engel (1998), with appropriate cardinal temperatures for each phase as reported by Setiyono *et al.* (2007). We did not account for photoperiod in our DS calculation as the four locations were located within a narrow latitudinal band (from 40.5°N to 42.6°N).

Seasonal dynamics of ADM and accumulated N were assessed in each treatment-replicate by collecting all plants within a 1-m section of row (inclusive of the 10 plants from which phenological development was assessed) surrounded by two rows receiving the same N treatment in order to avoid edge effects. We did not attempt to collect root biomass, which accounts for only *ca*. 10% of total plant biomass at R7 (Setiyono *et al.*, 2010b). Samples were collected weekly from VE until R7 and separated into green leaves, stems, seed, pod walls and senesced leaves. Green leaves (defined as any leaf with >50% green area) were scanned to determine the LAI in each ADM sample (LAI-3100 area meter LI-COR, Lincoln, NE). Abscised leaves were also collected every week within a 1-m long net placed between rows in each treatment-replicate. All samples were oven-dried at 70°C until reaching constant weight. Total ADM was calculated as the dry matter sum of all plant organs (included collected abscised leaves). Each plant organ sample was separately ground in a Wiley mill (1-mm screen mesh), and N concentration was determined with a dry combustion-based analyzer (LECO Corporation, St Joseph, MI). Larger plant samples (4.6 m<sup>2</sup>) were collected shortly after R7 from the two central rows in each plot, surrounded by two rows receiving the same N treatment, to obtain an end-of-season post-R7 estimate of seed yield. Two sub-samples of 200 seeds from the threshed seed of each plot were weighed to estimate the mean individual seed dry mass and to derive the number of seeds per harvested area. Seed yield and seed mass were adjusted to 130 g H<sub>2</sub>O kg<sup>-1</sup> seed, which is the market-based standard for soybean seed moisture content.

Seasonal dynamics of BNF were determined using the natural <sup>15</sup>N abundance method (Shearer & Kohl, 1986) in the zero-N treatment. Due to logistic constraints, BNF was measured in five of the seven environments. We present here data on BNF in the zero-N treatment only; determination of BNF in the full-N treatment was not possible due to the high isotopic fractionation as a result of increased denitrification with fertilizer addition (Mathieu *et al.*, 2007). The natural <sup>15</sup>N abundance method requires a 'reference crop' that does not fix N<sub>2</sub> and that is grown with the same N management of the legume crop (Peoples, Unkovich, & Herridge, 2009; Unkovich *et al.*, 2008). Following previous studies (Collino *et al.*, 2015; Pate, Unkovich, Armstrong, & Sanford, 1994), we used maize grown in 'N-omission' WILFY\_Plant. Ce

plots as the reference crop to measure BNF. A N-omission plot  $(9.1 \times 11 \text{ m})$  located adjacent (<30 m) to each soybean experiment was sown with maize on the same date as soybean and did not receive any N fertilizer (but received P and K fertilizer). Calculation of BNF following the natural <sup>15</sup>N abundance method requires that timing of N uptake between soybean and the reference crop is similar, regardless of the total amount of N absorbed (Witty, 1983). In our case, the maize plot was sown (and emerged) on the same date as soybean; maize relative maturity was purposely selected so that both maize and soybean crops reached physiological maturity around the same date. Also, rate of root depth over time and distribution of root length with depth are similar between the two crops, as has been documented in field studies conducted in the US Corn Belt (Nichols et al., 2019: Ordóñez et al., 2018). To summarize, we believe that our experimental setup was appropriate to ensure that the timing of N uptake coincided between the reference maize crop and soybean as it was required to calculate BNF following the natural <sup>15</sup>N abundance method.

Every week, we collected aboveground plant samples consisting of (a) 0.5-m row of consecutive soybean plants in the zero-N treatment, and (b) two maize plants in the N-omission maize plot. These samples were collected on the same dates as other plant samples for ADM and accumulated N determination. The natural <sup>15</sup>N abundance was measured using an elemental analyzer interfaced to a continuous flow isotope ratio mass spectrometer (UC Davis, California, USA). The fraction of the total N in ADM derived from BNF was calculated as:

$$\mathsf{BNF} = \frac{\delta 15 \mathsf{Nref} - \delta 15 \mathsf{Nsoy}}{\delta 15 \mathsf{Nref} - B} \tag{1}$$

where  $\delta^{15}$ Nref and  $\delta^{15}$ Nsov are the natural  $^{15}$ N abundance of the reference crop (maize grown in the N-omission plot) and soybean, respectively, and B is the natural <sup>15</sup>N abundance in soybean that relies only on BNF. The B value used in this work was -1.75, which corresponds to the average of the B values reported in the literature (Balboa & Ciampitti, 2020; Unkovich et al., 2008). Sensitivity analysis revealed that ±20% variation in the B value would have resulted in a range of BNF that fall within the experimental error, which justifies the use of an average B value from the literature. In the case of  $\delta^{15}$ Nref, we adjusted a quadratic model to smooth the observed variation among sampling times, with  $r^2$  of fitted models ranging from 0.67 to 0.93 across experiments (Supplementary Figure S2). The decline in  $\delta^{15}$ Nref over time was likely associated with the root system exploring soil layers with different <sup>15</sup>N enrichment as the crop season advances (Högberg, 1997; Shearer & Kohl, 1986). For a given sampling time, BNF (in kg N ha<sup>-1</sup>) was calculated based on the fraction of BNF and accumulated N; indigenous soil N supply was calculated as the difference between accumulated N and BNF. We do not expect results from our analysis to be influenced by the method selected for BNF determination as average BNF at R7 stage measured using the natural <sup>15</sup>N abundance method was identical to that estimated independently using the 'difference method' (i.e., absorbed N in soybean minus absorbed N in the reference maize crop). However, we reported here only the results on BNF derived from the natural <sup>15</sup>N abundance technique as it has been reported in the literature to be less prone to errors compared with the difference method (Unkovich *et al.*, 2008).

Incident and absorbed photosynthetically active radiation (PAR and APAR, respectively) were measured in one or two replicates in each treatment and environment. Measurements were taken every second and recorded as a 30-minute average, starting soon after VE and ending at R7. All sensors were calibrated by the manufacturer and a cross-calibration among sensors was performed every year before placing them in the field and again after removal at harvest. Sensors were leveled (if needed) and cleaned every 3 to 5 days during the season. The PAR was measured above the canopy using a point quantum sensor facing up (LI-190SA, LI-COR, Lincoln, NE). Transmitted PAR was measured with a singleline quantum sensor (LI-191SA, LI-COR, Lincoln, NE) placed at the soil surface diagonally across rows. Total (canopy plus soil) reflected PAR was measured using an inverted point quantum sensor (ibid) placed 2 m above crop canopy. Reflected PAR from soil was measured using an inverted line quantum sensor (ibid) placed 5 cm above soil surface and diagonally across rows. Daily canopy reflected PAR was calculated as total reflected PAR minus soil reflected PAR. Daily APAR was calculated as incident PAR minus transmitted PAR and canopy reflected PAR; and expressed as the fraction of daily incident PAR (fAPAR). Finally, radiation-use efficiency (RUE) was estimated as the slope of the relationship between accumulated ADM sampling points and their corresponding accumulated APAR from VE to R7 stages.

Leaf photosynthesis was measured at four stages during the growing season (R1, R3, R5 and R6) in all treatments and environments in 2017. All photosynthesis measurements were performed between 10 a.m. and 2 p.m., using only plants that were purposely selected based on any given above-stated crop stage so that all measured plants were at the same average crop stage recorded on that day. Photosynthesis was measured on the central leaflet of the third most recently developed leaf of one plant in each treatment-replicate. Light response curves were generated by varying the photosynthetic photon flux density (PPFD), from 1800 to 0  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> of a red/blue LED light source of an open-flow gas exchange system (LI-6400, Lincoln, NE). Carbon dioxide (CO<sub>2</sub>) concentration inside the chamber of the open-flow gas exchange system was maintained at 400 ppm, leaf temperature was set at 25°C, and the minimum measurement waiting time was 60 s or until reaching a coefficient of variation (CV) ≤ 3% of the CO<sub>2</sub> assimilation rate.

#### 2.4 | Data analysis

A logistic model (France & Thornley, 1984) was fitted to the weekly data of ADM and accumulated N collected from VE to R7:

Accumulated ADM or N = 
$$\frac{W0 * Wf}{W0 + (Wf - W0) * e^{-u * t}}$$
(2)

where W0 is the estimated ADM (g m<sup>-2</sup>) or N (kg ha<sup>-1</sup>) at emergence, Wf is the maximum ADM or accumulated N during the growing season, t is time in days after emergence, and u is a constant of

				Seed							
Location	Year	Treatment	Seed yield (Mg ha <sup>-1</sup> )	number (m <sup>-2</sup> )	Seed mass (mg)	ADM (Mg ha <sup>-1</sup> )	Accumulated N (kg N ha <sup>-1</sup> )	Seed N concentration (g N kg $^{-1}$ )	Mobilized ADM (Mg ha <sup>-1</sup> )	Mobilized N (kg N ha <sup>-1</sup> )	BNF (kg N ha <sup>−1</sup> ) <sup>∉</sup>
Mead	2016	Zero	5.3	3,320	161	12.8	419	56.8	1.0	154	264
		Full	6.2	3,606	172	13.7	460	57.1	1.5	174	n.c.
	2017	Zero	5.4	2,992	179	11.3	370	56.7	1.9	147	242
		Full	6.7	3,557	187	13.1	459	60.0	2.5	179	n.c.
Saronville	2016	Zero	5.8	3,405	171	12.1	408	57.7	0.9	134	270
		Full	6.4	3,476	184	13.4	436	58.0	1.0	140	n.c.
	2017	Zero	5.7	3,317	173	13.1	401	58.0	2.1	167	n.c.
		Full	6.4	3,584	179	14.7	485	62.7	1.6	196	n.c.
Smithfield	2016	Zero	5.3	3,031	174	10.4	364	54.7	1.7	130	293
		Full	5.7	3,065	186	11.5	384	57.5	1.1	157	n.c.
	2017	Zero	5.5	2,676	204	11.2	367	62.3	1.5	151	218
		Full	5.5	2,813	197	12.4	432	64.5	1.0	172	n.c.
Atkinson	2017	Zero	5.5	2,933	188	13.1	436	61.2	1.5	165	n.c.
		Full	5.7	3,009	191	13.3	466	61.6	1.8	214	n.c.
<i>Note</i> : Also sho	wn is bic	ological N fixati	ion (BNF) measure	d for the zero N	treatment.						

<sup>a</sup>BNF measured for a subset of five environments. BNF was not calculated (n.c.) for the full-N treatment given the high isotopic fractionation due to the large amount of N applied (see Section 2.3).

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		Seed yield	Seed number	Seed mass	ADM	Accumulated N	Seed N concentration	Mobilized ADM	Mobilized N
Fixed effects	d.f.	F	F	F	F	F	F	F	F
E	6	7.3***	20.4***	59.2***	16.1***	5.9**	4.6**	5.3**	5.0**
E	6	7.3***	20.4***	59.2***	16.1***	5.9**	4.6**	5.3**	5.0**
Ν	1	42.6***	18.7***	24.8***	16.8***	27.7***	5.6*	0.2	16.8***
N x E	6	3.0*	2.2	4.1**	0.5*	1.2	0.3	2.9*	0.6
Random effects		MS	MS	MS	MS	MS	MS	MS	MS
Rep (E)	21	0.08	31,207	15.7	0.5	1,011	11	0.25	566
N x rep (E)	21	0.12	31,437	24.1	1.3	1,326	10	0.17	578
Estimated means		(Mg ha <sup>-1</sup> )	(m <sup>-2</sup> )	(mg)	(Mg ha <sup>-1</sup> )	(kg N ha <sup>-1</sup> )	(g N kg ADM <sup>-1</sup> )	(Mg ha <sup>-1</sup> )	(kg N ha $^{-1}$ )
Full-N		6.1	3,301	185	13.2	446	60	1.5	176
Zero-N		5.5	3,096	179	12.0	395	58	1.5	150
Difference (full-N minus zero-N)		0.6***	205***	7***	1.2***	51***	2*	Nil	26***

**TABLE 3** Analysis of variance for effect of nitrogen (N) treatment on soybean seed yield, number and mass, aboveground dry matter (ADM), accumulated nitrogen (N) in ADM, and seed N concentration at physiological maturity (R7 stage), and mobilized ADM and N from non-seed ADM.

*Note*: Each experiment was considered to be a separate environment (E) for the N  $\times$  E interaction in this analysis. Also shown are the F-test values and probabilities for the fixed effects, mean squares (MS) for the random effects, estimated means for each treatment, and contrast between N treatments means. Asterisks indicate statistical significance at \**P* < 0.01, and \*\*\**P* < 0.001.

proportionality of plant growth and its decaying with time. The derivative of Equation 2, with respect to time, represents the crop growth rate (CGR; g m<sup>-2</sup> d<sup>-1</sup>) or N accumulation rate (NA<sub>rate</sub>; kg N ha<sup>-1</sup> d<sup>-1</sup>). Estimating CGR and NA<sub>rate</sub> using this approach helps remove the measurement error associated with specific sampling times (*e.g.*, Bange, Hammer, & Rickert, 1997; Hall, Connor, & Sadras, 1995; Lindquist *et al.*, 2005). Similarly, a Gaussian distribution model was fit to the seasonal dynamics of LAI:

$$LAI = LAImax * e^{\left(-0.5 * \left(\frac{t-tLAImax}{SD}\right)^{2}\right)}.$$
 (3)

where LAImax is estimated maximum LAI,  $t_{LAImax}$  is time at which at LAImax is reached, and SD is a parameter of the exponential equation.

Seasonal dynamics in fAPAR were compared between N treatments. Because we did not have light sensors in every plot, we used the following approach to estimate daily fAPAR for each experimental unit. First, we generated a relationship between measured fAPAR and LAI using all available dates of LAI sampling (Supplementary Figure S3). The extinction coefficient (*k*) was estimated to be 0.54 across environments and treatments, without differences between N treatments. Second, we used that relationship to derive the daily fAPAR for each treatment in each experiment based on daily LAI obtained from the fitted models (Equation 3).

Sources of carbon for seed dry matter accumulation during the reproductive phase include new photo-assimilates created on a daily basis (Yamagata, Kouchi, & Yoneyama, 1987), and dry matter mobilization from vegetative organs (Egli, Guffy, & Leggett, 1985; Stephenson & Wilson, 1977). Apparent dry matter mobilization to seeds was estimated as the difference between non-seed ADM (including stems, green,



**FIGURE 2** Observed seed number and seed mass at physiological maturity in the full-N (red squares) and zero-N (blue circles) treatments across the seven environments. Connecting grey lines relate paired treatments from the same environment. Dotted lines indicate different seed yields across the seed number and mass ranges. Parameters of the fitted linear regressions (solid blue and red lines for zero-N and full-N treatment, respectively) and coefficients of determination ( $r^2$ ) are also shown. Note that regression lines do not imply causality (with respect to one trait *versus* the other); instead, they are shown to illustrate the trade-off between seed number and mass for each N treatment found across the seven irrigated experiments in Nebraska

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senesced and abscised leaves, and pod walls) at R5 and R7. We used the term 'apparent' because the amount of non-seed ADM mobilized to the seeds was calculated as the difference in non-seed ADM between R5 and R7, rather than being directly measured, and also to recognize that it does not account for the conversion efficiency (Borrás, Slafer, & Otegui, 2004). In the case of seed N accumulation, major sources include crop soil N uptake and BNF during seed filling as well as mobilization from non-seed ADM to the seeds. This mobilized N was estimated following the same approach as for dry matter mobilization. We did not attempt to account for N mobilized from belowground biomass, which we estimated to represent a very low fraction (*ca.* 3%) of the total N mobilized to seed, based on root-to-shoot ratio, root N concentration, and N mobilization fraction reported in the literature (Amthor *et al.*, 1994; Connor, Loomis, & Cassman, 2011; Setiyono *et al.*, 2010b).

Following Connor *et al.* (2011), the following function was used to quantify the response of net photosynthesis to incident light:

$$A = \frac{Amax * (PPFD - Ic) * \alpha}{Amax + (PPFD - Ic) * \alpha}$$
(4)

where A is the photosynthetic rate ( $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>), Amax is the maximum photosynthetic rate at light saturation ( $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>), PPFD is photosynthetic photon flux density ( $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>), *lc* is the light compensation point ( $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>) and  $\alpha$  is the initial slope of the response curve. Dark respiration (*Rd*;  $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>) can be estimated from the fitted parameter values as  $-\alpha^* lc$ .

We used a combined analysis of variance (ANOVA) for the effect of N treatments across environments where environments and N treatments were treated as fixed effects (SAS<sup>®</sup> PROC MIXED v.9.3; Moore & Dixon, 2015). The combined ANOVA assumes homogeneity of the error variances within a given environment and when pooled over all seven environments. Our analysis indicated that the error variances did not deviate much from homogeneity ( $F_{max}$  <6), which led us to consider our analysis to be robust (Milliken & Johnson, 2009). The following directly observed or calculated parameters were evaluated: seed yield, seed number, seed mass, ADM at R7, accumulated N at R7, seed N concentration and mobilized ADM and N. A similar analysis was performed to identify the crop stage interval with the largest

**FIGURE 3** (a) Soybean aboveground dry matter (ADM), and (b) nitrogen (N) accumulation in the full-N (red squares) and zero-N (blue circles) treatments as a function of development stage (DS) on the bottom x-axis. (c) Same as (b) but also showing accumulated N derived from indigenous soil N (ISN; brown downward triangles) and biological N fixation (BNF; green upward triangles) in the zero-N treatment. Vn and Rn stages based on Fehr & Caviness (1977) are shown in the top x-axis. Solid red and blue lines represent the fitted Equation 2 for the full-N and zero-N treatment means computed from pooled data across environments. Insets show sigmoid derivative daily crop growth rates (CGR), N accumulation rates (NA<sub>rate</sub>), and rates of ISN and BNF. Coefficient of determination of fitted models was >0.90 in all cases [Colour figure can be viewed at wileyonlinelibrary.com]

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**FIGURE 4** (a) Relationship between seed number and accumulated aboveground dry matter (ADM) between the beginning of pod setting (R3) to full seed (R6) stages in the full-N (red squares) and zero-N (blue circles) treatments. Connecting lines relate paired treatments from the same environment. Parameters of the fitted linear regression (solid black line) and coefficient of determination ( $r^2$ ) are shown. (b) Comparison of accumulated ADM between the R3-R6 phase between the full-N and zero-N treatments for the seven environments. Solid black line labeled 1:1 indicates *y* = *x* null hypothesis of no difference [Colour figure can be viewed at wileyonlinelibrary.com]

900 1000 1100

difference between the two N treatments relative to CGR and NA<sub>rate</sub>. To do this, the CGR and NA<sub>rate</sub> obtained per replicate were averaged by treatment over four crop phases (VE-R1, R1-R3, R3-R5, R5-R7). Variation in seed yield is largely driven by differences in seed set, which, in turn, is modulated by the crop growth rate during key reproductive phases; the latter has been referred to as 'critical period' (Egli & Bruening, 2006; Fischer, 1975). Following Kantolic, Peralta, & Slafer (2013), accumulated ADM during the critical period for soybean (between R3 and R6 stages) was used to understand seed number differences between N treatments. Least square means were computed for each treatment, and treatment differences evaluated using Dunnett's test.

800

ADM R3-R6 (g m<sup>-2</sup>)

700

600

Differences in the parameters of the light response curve (*Amax*, *lc*,  $\alpha$ , *Rd*) between N treatments were evaluated using mixed models (InfoStat; Di Rienzo *et al.*, 2011). Linear regression analysis was used to investigate relationships and trade-offs between the measured and calculated variables. Slope and intercept were calculated and their differences between treatments were tested with *F* tests. To remove the confounding effect of differences in phenology across environments, seasonal patterns in ADM, accumulated N, CGR, NA<sub>rate</sub> and LAI are shown as a function of the DS calculated for each environment (Figure 1), but with data across environments pooled for each N treatment to facilitate the comparison.

#### 3 | RESULTS

## 3.1 | Impact of soybean nitrogen limitation on seed yield components

Soybean seed yield ranged from 5.3 to 5.8 Mg  $ha^{-1}$  under zero N and from 5.5 to 6.7 Mg  $ha^{-1}$  with non-limiting N across the seven

environments (Table 2). In all cases, measured yields in the full-N treatment were within 15% of the simulated yield potential (Table 1). Average seed yield was 0.6 Mg ha<sup>-1</sup> greater in the full-N than in the zero-N treatment (Table 3). Total accumulated N was 51 kg N ha<sup>-1</sup> greater in the full-N than in the zero-N treatment (13%; 446 versus 395 kg N ha<sup>-1</sup>, respectively), which translated to 10% greater ADM at R7 in the full-N compared with the zero-N treatment (13.2 versus 12.0 Mg ha<sup>-1</sup>, respectively). The full-N treatment exhibited 7% greater seed number (3.301 versus 3.096 seeds  $m^{-2}$ ), and 4% greater seed mass (185 versus 179 mg) than the zero-N treatment. Ordinarily, in absence of strong environmental variation, seed number and seed mass are negatively correlated, and that was the case across our irrigated soybean experiments in NE (Figure 2). However, the slope of the linear regression between seed mass and seed number was significantly different between the zero-N and full-N treatments (-0.02 versus -0.05 mg per additional seed, respectively; P = 0.005), revealing that the seed mass versus seed number trade-off was substantively alleviated in the case of the full-N treatment (Figure 2).

700

800

ADM<sub>zero-N</sub> R3-R6 (g m<sup>-2</sup>)

600

900

1000 1100

## 3.2 | Differences in seasonal accumulated ADM and N between treatments

Accumulated ADM and N followed sigmoidal patterns (Figure 3). The sigmoidal inflection points corresponded to *ca*. R5 and to a DS value of *ca*. 1.25, which are more precisely documented in the inset plots that show the peaks of the derivative variables CGR and NA<sub>rate</sub>. It was also evident in the sigmoid patterns that greater ADM and accumulated N became visibly different just before R1 in the full-N compared with the zero-N treatment. However, the post-peak downward slopes after R5 for CGR and NA<sub>rate</sub> were coincident between N treatments. Maximum CGR was slightly higher in the full-N than in the zero-N





**FIGURE 5** (a) Soybean leaf area index and (b) estimated fraction of absorbed photosynthetically active radiation (fAPAR) in the full-N (red squares) and zero-N (blue circles) treatments as a function of developmental stage (DS). Stages based on Fehr & Caviness (1977) are shown in the top x-axis. Solid lines represent the fitted models for the full-N (red) and zero-N treatment (blue) based on the pooled data across environments. Data for each N treatment were pooled across environments. See Section 2.4 for detailed explanation on fAPAR estimation [Colour figure can be viewed at wileyonlinelibrary.com]

treatment (22.4 versus 21.7 g m<sup>-2</sup> d<sup>-1</sup>; P = 0.046), but the growth stage at which the peak of CGR occurred did not differ between N treatments (P = 0.212; Figure 3a, inset). In contrast, maximum NA<sub>rate</sub> was similar between N treatments (7.7 versus 7.4 kg N ha<sup>-1</sup> d<sup>-1</sup> in the full-N and zero-N treatments, respectively; P = 0.180); however, the peak of NA<sub>rate</sub> occurred earlier in the full-N compared with the zero-N treatment (P = 0.001; Figure 3b, inset). On average, CGR and NA<sub>rate</sub> between VE-R5 were 11% and 22% higher in the full-N

compared with the zero-N treatment, respectively (Supplementary Table S1 and S2). These differences led to an additional 676 kg ADM  $ha^{-1}$  and 45 kg N  $ha^{-1}$  accumulated at R5 in the full-N *versus* zero-N treatment.

On average, indigenous soil N and BNF accounted for 33% and 67% of the accumulated N at R7 stage (Figure 3c, Table 2). However, relative contribution of indigenous soil N and BNF to plant N accumulation varied during the season. For example, indigenous soil N accounted for the largest portion of accumulated N between VE and R5 stages (65%), while BNF supplied most of the N during the seed filling (90%). Maximum rates of N accumulation from indigenous soil N and BNF occurred around R3 (DS = 1.07) and R5.5 stages (DS = 1.56), respectively (Figure 3c, inset).

Accumulated N in the full-N treatment can be taken as a measure of plant N demand when N supply is not limiting. Comparison of rates of accumulated N in full-N treatment *versus* BNF in the zero-N treatment is of interest as to discern when (and the extent to which) the combined N supply from BNF and indigenous soil N was not sufficient to meet plant N demand. Daily rates of plant N demand and BNF increased gradually during the season until reaching a peak, declining subsequently during the seed filling (Figure 3c, inset). However, the peak of BNF rate occurred later compared with plant N demand (DS = 1.54 *versus* 1.28) and daily BNF was consistently lower than the plant N demand during the entire crop cycle until the middle of the seed filling phase. During the same period, NA<sub>rate</sub> was consistently lower in the zero-N than in the full-N treatment, suggesting that BNF was not sufficient to fill in the 'N gap' between plant N demand and indigenous soil N supply.

# 3.3 | Drivers for differences in seed number between N treatments

Seed number was positively associated with accumulated ADM between the R3 and R6 stages (Figure 4a). No differences in slope or intercept of the relationship between seed number and accumulated ADM during R3-R6 period were detected between the two N treatments (P > 0.60), so N treatment data were pooled. Inferentially, this means that number of seeds set per unit of accumulated ADM between R3 and R6 remained unchanged between N treatments. Accumulated ADM between R3 and R6 was greater in the full-N than in the zero-N treatment (816 *versus* 785 g m<sup>-2</sup>, respectively; paired *T*-test, P = 0.045) (Figure 4b), leading to higher seed number in the full-N compared with the zero-N treatment (Tables 2 and 3). Differences in R3-R6 duration between N treatments was not significant (P = 0.356), indicating that differences in accumulated ADM during the R3-R6 phase between N treatments was associated with differences in CGR (Supplementary Table S1, and S2).

## 3.4 | Mechanisms explaining differences in accumulated ADM between N treatments

The two N treatments differed in terms of their impact on the seasonal dynamics of LAI (Figure 5a). First, LAI early in the season was greater in

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**FIGURE 6** Aboveground biomass plotted against cumulative absorbed photosynthetically active radiation (APAR) in the full-N (red squares) and zero-N (blue circles) treatments across the seven environments. The upper left panel shows the pooled data from the seven environments. Slope of the fitted linear regression represents the radiation-use efficiency (RUE; g  $MJ^{-1}$ ). Significance of the statistical test for the null hypothesis of no difference between slopes between N treatments is shown. Asterisks indicate significance at \**P* < 0.10, \*\**P* < 0.05, or \*\*\**P* < 0.001 [Colour figure can be viewed at wileyonlinelibrary.com]

the full-N than in the zero-N treatment, which led to greater leaf area duration before R5 (*i.e.*, the integral of the LAI curve before R5; P = 0.010). Second, while the maximum LAI value was nearly identical in the full-N and zero-N treatments (5.8 and 5.7, respectively; P = 0.804), the LAImax value was reached sooner in the full-N compared with the zero-N treatment (1.22 *versus* 1.37 DS units, respectively; P = 0.007). Third, the rate of (post-max) decline in LAI values was greater in the full-N than in the zero-N treatment, though LAI trends of the two N treatments converged towards the end of the season.

Seasonal patterns in estimated fAPAR followed the observed dynamics in LAI (Figure 5b). The full-N treatment exhibited consistently greater fAPAR between VE and R5, reaching *ca.* 95% of full interception 3 days earlier (P < 0.001) compared with the zero-N treatment. These differences resulted in larger total estimated cumulative APAR from VE to R7 stages in the full-N than in the zero-N treatment (700 *versus* 679 MJ m<sup>-2</sup>; P = 0.014).

Canopy-level radiation-use efficiency was significantly greater in the full-N compared with the zero-N treatment in six of the seven environments (Figure 6). The RUE was 8% greater in the full-N than in the zero-N treatment (2.00 *versus* 1.86 g MJ<sup>-1</sup>, respectively). In contrast, we could not detect significant differences in leaf-level photosynthesis (*i.e.*, net CO<sub>2</sub> assimilation rate) between N treatments (Figure 7). Crop stage (*i.e.*, DS) significantly influenced *Amax* and *lc* (P < 0.04), with *Amax* increasing until reaching a maximum (40.8 µmol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>) at *ca*. R5 (DS = 1.3) stage, but with *lc* and *Rd* gradually decreasing from the first sampling date just prior to stage R1 onward

to the last sampling date at R6 stage. Changes in  $\alpha$  were small, without a clear pattern. Overall, the above four photosynthetic parameters, and also leaf N concentration and specific leaf weight (SLW), were not statistically significant between N treatments (P > 0.1), but the full-N treatment tended to have greater leaf N concentration and SLW compared with the zero-N treatment (Supplementary Figure S4).

## 3.5 | Drivers for differences in seed mass and seed N between N treatments

Rate of seed dry matter accumulation during the seed filling period (R5-R7 phase) was greater in full-N than in the zero-N treatment (P = 0.032). There was a strong relationship between mobilized N from non-seed ADM to seed and the amount of N in non-seed ADM at R5 stage (Figure 8a). Differences in mobilized N between the treatments were associated with greater accumulated N at R5 in the full-N compared with the zero-N treatment (281 versus 242 kg N ha<sup>-1</sup>, respectively; P < 0.001), without changes in the fraction of non-seed N that was mobilized between N treatments (62%; P = 0.525). Indeed, mobilized N was 17% greater in the full-N than in the zero-N treatment (176 versus 150 kg N ha<sup>-1</sup>, respectively, P = 0.001; Table 3, Figure 8b). Apparent dry matter mobilization from non-seed ADM to seed was smaller than N mobilization (*ca.* 14% of non-seed ADM at R5) and not affected by the N treatments (P = 0.283; Table 3).



**FIGURE 7** Leaf net photosynthesis (A) as a function of photosynthetic photon flux density (PPFD) in the full-N (red squares) and zero-N (blue circles) treatments at four development stages (DS). Parameters of the fitted models are shown: maximum photosynthesis in  $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup> (*Amax*), light compensation point in  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> (*Ic*), initial slope of light response curve ( $\alpha$ ), and dark respiration in  $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>(*Rd*). Data for each N treatment were pooled across environments [Colour figure can be viewed at wileyonlinelibrary.com]

### 4 | DISCUSSION

In highly productive soybean systems, the combined N supply from BNF and indigenous soil N supply cannot meet the crop N requirement as indicated by the differences in  $NA_{rate}$  between the N treatments before R5 stage (Figure 3b). There was an asynchrony between BNF and plant N demand: BNF lagged behind plant N demand (i.e., started and peaked later) and it was not sufficient to meet crop N demand after accounting for the indigenous soil N supply (Figure 3c). Sequential N fertilizer application induced a greater  $NA_{rate}$ , as indicated by the difference between the full-N and zero-N treatments, that in turn supported faster leaf area expansion which, combined with greater fAPAR and CGR, allowed a larger seed set during the R3-R6 phase (Figures 3, 4, and 5; Supplementary Table S1 and S2). Similarly, the extra N accumulated in the ADM before R5 in the full-N compared with the zero-N treatment (+45 kg N ha<sup>-1</sup>) resulted in greater N mobilization from non-seed ADM to seed (+26 kg N  $ha^{-1}$ ) (Figure 8, Tables 2 and 3). This is consistent with Sinclair, Farias, Neumaier, and Nepomuceno (2003) and Sinclair & Rufty (2012), who postulated that the degree to which the plant can supply N to meet seed requirements will determine its capacity to meet the potential seed filling rate. These results also suggest that strategies to increase N supply in soybean should aim to increase NA<sub>rate</sub> before seed filling, with the goal of increasing the CGR during the critical period for seed number determination, and increase the amount of N in non-seed ADM at R5 to support a greater seed mass and seed N concentration. Finally, leaf-level photosynthesis did not differ among N treatments. In contrast, canopy-level RUE was greater in the full-N compared with the zero-N treatment, which might be associated with changes in root-to-shoot ratio, reduced costs due to lower BNF as a result of N fertilizer application, and/or variation in leaf N distribution within the canopy (Bonelli & Andrade, 2020; Cassman, Whitney, & Stockinger, 1980; Pate & Layzell, 1990).

The zero-N treatment seed yield varied within a narrow range (from 5.3 to 5.8 Mg ha<sup>-1</sup>) suggesting that high-yield soybean relying exclusively on N supply from soil and BNF has an upper yield limit near 5.5 Mg ha<sup>-1</sup> in Nebraska (Figure 2, Table 2). In contrast, in four of the seven environments, seed yield in the full-N treatment reached ca. 6.5 Mg ha<sup>-1</sup> because of a simultaneous increase in both seed number and mass. In the other three environments, the full-N treatment yields were lower (5.5-5.7 Mg ha<sup>-1</sup>), and thus closer to the zero-N treatment yields, which was attributable to the high indigenous soil N supply (ca. 100-150 kg N ha<sup>-1</sup>) at these three site-years as documented previously in Cafaro La Menza et al. (2019) together with the relatively lower yield potential at these three site-years (Table 1). Similarly, it was remarkable also to find a simultaneous increase in seed yield and seed N concentration given the well-documented trade-off between these two variables (Chung et al., 2003). The degree to which seed N demand is met in soybean depends on (a) NA<sub>rate</sub> during seed filling (R5-R7), which includes N from soil and BNF, and (b) N mobilized from non-seed ADM to the growing seeds (Egli et al., 1985; Stephenson & Wilson, 1977; Yamagata et al., 1987). In our study, the NA<sub>rate</sub> differed between N treatments, except for the R5-R7 seed



**FIGURE 8** (a) Relationship between N mobilized from non-seed dry matter to seed during seed filling and the amount of accumulated N in non-seed dry matter at R5 in the full-N (red squares) and zero-N (blue circles) treatments. Parameters of the fitted linear regression (solid black line) and coefficient of determination ( $r^2$ ) are shown. (b) Comparison of mobilized N between the full-N and zero-N treatments. Solid black line labeled 1:1 indicates y = x null hypothesis of no difference. Parameters of the fitted linear regression (dashed line) and coefficient of determination ( $r^2$ ) are also shown [Colour figure can be viewed at wileyonlinelibrary.com]

filling phase. In turn, the amount of mobilized N was greater in the full-N compared with the zero-N treatment as a result of higher accumulated N in the non-seed ADM at R5, with no change in the fraction of N mobilized compared with the zero-N treatment. Clearly, the extra accumulated N in non-seed ADM at R5 in the full-N versus zero-N treatment should not be seen as 'luxury N consumption' as it helped increase seed mass and maintain seed N concentration (Staswick, 1994). Even when soil N was non-limiting in the full-N treatment, greater remobilized N (rather than NA<sub>rate</sub>) was the underlying mechanism explaining greater seed mass and seed N concentration compared with the zero-N treatment. These results are consistent with the CROPGRO model (Boote, Jones, Hoogenboom, & Pickering, 1998) in which mobilized N from non-seed organs to seed is calculated as a function of thermal time and unaffected by soil N availability during the seed filling. This preference for remobilizing N from non-seed ADM may be associated with the lower cost of protein breakdown and re-synthesis compared with de novo protein synthesis from indigenous soil N supply and BNF (De Vries, Van Laar, & Chardon, 1983). It may also indicate a loss of root functionality during seed filling as has been reported for other crop species (Lisanti, Hall, & Chimenti, 2013; Thibodeau & Jaworski, 1975). Understanding the underlying drivers behind the apparent limited capacity of the plant to make use of available soil N during seed filling deserves further research.

In a global context, crop yields need to increase *ca*. 50% by year 2050 in order to meet food demand and avoid a massive conversion of natural ecosystems into cropland (Cassman & Grassini, 2020). For soybean, the important issue is the degree to which N limitation may or not allow such a yield achievement. Given the current average yield of *ca*. 3 Mg ha<sup>-1</sup> in major producing areas, the goal would have to be an average field yield of 4.5 Mg ha<sup>-1</sup> by year 2050, which entails a N requirement of 360 kg N ha<sup>-1</sup>. Indigenous soil N supply may be able to cover half of that N requirement considering that fertile agricultural

soils in USA and Argentina can provide *ca*. 100–150 kg N ha<sup>-1</sup> (Cafaro La Menza et al., 2019). The remaining N requirement (ca. 200-250 kg N ha<sup>-1</sup>) can be readily supplied by BNF (Salvagiotti et al., 2008). Thus, adoption of improved agronomic practices and cultivars can be cost-effective options to increase yields in these intermediate-yield production systems (e.g., Rattalino Edreira et al., 2017; Di Mauro et al., 2018). However, much higher BNF would be required to sustainably advance yield gains in high-yield irrigated soybean production areas such as Nebraska and other areas in the Central US Great Plains where producers are already achieving average yields of ca. 4.5 Mg ha<sup>-1</sup>. In these environments, a 50%-yield increase would imply an average yield of 6.8 Mg ha<sup>-1</sup> by year 2050, which has an associated N requirement of 540 kg N ha<sup>-1</sup>. Assuming the same level of indigenous soil N (150 kg N ha<sup>-1</sup>), BNF would then have to increase to 400 kg N ha<sup>-1</sup>. None of the recent reviews on BNF on soybean provide evidence that reaching such a high level of BNF and indigenous soil N supply is possible (Ciampitti & Salvagiotti, 2018; Salvagiotti et al., 2008). Our N fertilizer treatment was successful at increasing both yield and seed N concentration. However, it was far from being cost-effective and obviously not an environmentally sound practice to be adopted in commercial farms. Increasing BNF and/or indigenous soil N supply, improving the synchrony between N fixation and plant N demand, and alleviating the trade-off between the two sources of N are avenues worth exploring, even though the associated probability of success and timeline for impact are unknown (Denison & Kiers, 2005; Giller & Cadisch, 1995; Van Kessel & Hartley, 2000).

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#### **CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest regarding the publication of this article.

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#### REFERENCES

- Amthor, J. S., Mitchell, R. J., Runion, G. B., Rogers, H. H., Prior, S. A., & Wood, C. W. (1994). Energy content, construction cost and phytomass accumulation of *Glycine max* (L.) Merr. and *Sorghum bicolor* (L.) Moench grown in elevated CO<sub>2</sub> in the field. *New Phytologist*, 128, 443–450.
- Balboa, G. R., & Ciampitti, I. A. (2020). Estimating biological nitrogen fixation in field-grown soybeans: Impact of B value. *Plant and Soil*, 446, 195–210.
- Bange, M. P., Hammer, G. L., & Rickert, K. G. (1997). Environmental control of potential yield of sunflower in the subtropics. Australian Journal of Agricultural Research, 48, 231–240.
- Barraclough, P. B., Howarth, J. R., Jones, J., Lopez-Bellido, R., Parmar, S., Shepherd, C. E., & Hawkesford, M. J. (2010). Nitrogen efficiency of wheat: Genotypic and environmental variation and prospects for improvement. *European Journal of Agronomy*, 33, 1–11.
- Bender, R. R., Haegele, J. W., & Below, F. E. (2015). Nutrient uptake, partitioning, and remobilization in modern soybean varieties. Agronomy Journal, 107, 563–573.
- Bonelli, L. E., & Andrade, F. H. (2020). Maize radiation use-efficiency response to optimally distributed foliar-nitrogen-content depends on canopy leaf-area index. *Field Crops Research*, 247, 107557.
- Boote, K. J., Jones, J. W., Hoogenboom, G., & Pickering, N. B. (1998). Simulation of crop growth: CROPGRO model. In R. M. Peart & W. D. Shoup (Eds.), Agricultural systems modeling and simulation (pp. 651–692). New York, NY: Marcel Dekker Inc.
- Borrás, L., Slafer, G. A., & Otegui, M. E. (2004). Seed dry weight response to source-sink manipulations in wheat, maize and soybean: A quantitative reappraisal. *Field Crops Research*, *86*, 131–146.
- Brumm, T. J., & Hurburgh, C. R., Jr. (2006). Changes in long-term soybean compositional patterns. *Journal of the American Oil Chemists' Society*, 83, 981–983.
- Cafaro La Menza, N., Monzon, J. P., Specht, J. E., & Grassini, P. (2017). Is soybean yield limited by nitrogen supply? *Field Crops Research*, 213, 204-212.
- Cafaro La Menza, N., Monzon, J. P., Specht, J. E., Lindquist, J. L., Arkebauer, T. J., Graef, G., & Grassini, P. (2019). Nitrogen limitation in high-yield soybean: Seed yield, N accumulation, and N-use efficiency. *Field Crops Research*, 237, 74–81.
- Carciochi, W. D., Rosso, L. H. M., Secchi, M. A., Torres, A. R., Naeve, S., Casteel, S. N., ... Ciampitti, I. A. (2019). Soybean yield, biological N<sub>2</sub> fixation and seed composition responses to additional inoculation in the United States. *Scientific Reports*, 9, 1–10.
- Cassman, K. G., & Grassini, P. (2020). A global perspective on sustainable intensification research. *Nature Sustainability*, 3, 262–268.
- Cassman, K. G., Whitney, A. S., & Stockinger, K. R. (1980). Root growth and dry matter distribution of soybean as affected by phosphorus stress, nodulation, and nitrogen source. *Crop Science*, 20, 239–244.
- Chung, J., Babka, H. L., Graef, G. L., Staswick, P. E., Lee, D. J., Cregan, P. B., ... Specht, J. E. (2003). The seed protein, oil, and yield QTL on soybean linkage group I. Crop Science, 43, 1053–1067.

- Ciampitti, I. A., & Salvagiotti, F. (2018). New insights into soybean biological nitrogen fixation. Agronomy Journal, 110, 1185–1196.
- Collino, D. J., Salvagiotti, F., Perticari, A., Piccinetti, C., Ovando, G., Urquiaga, S., & Racca, R. W. (2015). Biological nitrogen fixation in soybean in Argentina: Relationships with crop, soil, and meteorological factors. *Plant and Soil*, 392, 239–252.
- Connor, D. J., Loomis, R. S., & Cassman, K. G. (2011). Crop ecology: Productivity and management in agricultural systems (2nd ed.) Cambridge: Cambridge University Press.
- De Bruin, J. L., & Pedersen, P. (2009). New and old soybean cultivar responses to plant density and intercepted light. *Crop Science*, 49, 2225–2232.
- De Bruin, J. L., Pedersen, P., Conley, S. P., Gaska, J. M., Naeve, S. L., Kurle, J. E., ... Abendroth, L. J. (2010). Probability of yield response to inoculants in fields with a history of soybean. *Crop Science*, 50, 265–272.
- De Vries, F. P., Van Laar, H. H., & Chardon, M. C. M. (1983). Bioenergetics of growth of seeds, fruits and storage organs. In S. J. Banta (Ed.), *Potential* productivity of field crops under different environments (pp. 37–60). Los Baños, Philippines: International Rice Research Institute.
- Denison, R. F., & Kiers, E. T. (2005). Sustainable crop nutrition: Constraints and opportunities. In M. Broadley (Ed.), *Plant nutritional genomics* (pp. 242–264). Oxford, UK: Blackwell Publishing.
- Di Rienzo, J. A., Casanoves, F., Balzarini, M. G., González, L., Tablada, M., & Robledo, Y. C. (2011). InfoStat v.2011. Universidad Nacional de Córdoba, Argentina. Retrieved from http://www.infostat.com.ar.
- Di Mauro, G., Cipriotti, P. A., & Rotundo, J. L. (2018). Environmental and management variables explain soybean yield gap variability in Central Argentina. *European Journal of Agronomy*, 99, 186–194.
- Egli, D. B., & Bruening, W. P. (2006). Temporal profiles of pod production and pod set in soybean. *European Journal of Agronomy*, 24, 11–18.
- Egli, D. B., Guffy, R. D., & Leggett, J. E. (1985). Partitioning of assimilate between vegetative and reproductive growth in soybean. Agronomy Journal, 77, 917–922.
- Farmaha, B. S., Eskridge, K. M., Cassman, K. G., Specht, J. E., Yang, H., & Grassini, P. (2016). Rotation impact on on-farm yield and input-use efficiency in high-yield irrigated maize-soybean systems. Agronomy Journal, 108, 2313–2321.
- Fehr, W. R., & Caviness, C. E. (1977). Stages of soybean development. Special Report 80. Iowa Agriculture and Home Economics Experiment Station, Iowa State University, Ames.
- Fischer, R. A. (1975). Yield potential in a dwarf spring wheat and the effect of shading. *Crop Science*, 15, 607–613.
- France, J., & Thornley, J. H. (1984). Mathematical models in agriculture. London, UK: Butterworths.
- Giller, K. E., & Cadisch, G. (1995). Future benefits from biological nitrogen fixation: An ecological approach to agriculture. *Plant and Soil*, 174, 255–277.
- Grassini, P., Specht, J., Tollenaar, T., Ciampitti, I., Cassman, K. G. (2014) Highyield maize-soybean cropping systems in the U.S. Corn Belt. In *Crop Physiology: applications for genetic improvement and agronomy* (V. O. Sadras Calderini, D. F.), 2nd), pp 17–41. Elsevier, The Netherlands.
- Hall, A. J., Connor, D. J., & Sadras, V. O. (1995). Radiation-use efficiency of sunflower crops: Effects of specific leaf nitrogen and ontogeny. *Field Crops Research*, 41, 65–77.
- Högberg, P. (1997). <sup>15</sup>N natural abundance in soil-plant systems. New Phytologist, 137, 179–203.
- Irmak, S., Payero, J. O., VanDeWalle, B., Rees, J., & Zoubek, G. (2014). Principles and operational characteristics of Watermark granular matrix sensor to measure soil water status and its practical applications for irrigation management in various soil textures. Extension Circular EC783. University of Nebraska-Lincoln. Retrieved from https://digitalcommons.unl.edu/biosysengfacpub/332/.
- Kantolic, A. G., Peralta, G. E., & Slafer, G. A. (2013). Seed number responses to extended photoperiod and shading during reproductive stages in indeterminate soybean. *European Journal of Agronomy*, 51, 91–100.

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- Leggett, M., Diaz-Zorita, M., Koivunen, M., Bowman, R., Pesek, R., Stevenson, C., & Leister, T. (2017). Soybean response to inoculation with *Bradyrhizobium japonicum* in the United States and Argentina. *Agronomy Journal*, 109, 1031–1038.
- Lindquist, J. L., Arkebauer, T. J., Walters, D. T., Cassman, K. G., & Dobermann, A. (2005). Maize radiation use efficiency under optimal growth conditions. *Agronomy Journal*, 97, 72–78.
- Lisanti, S., Hall, A. J., & Chimenti, C. A. (2013). Influence of water deficit and canopy senescence pattern on *Helianthus annuus* (L.) root functionality during the grain-filling phase. *Field Crops Research*, 154, 1–11.
- Mathieu, O., Lévêque, J., Hénault, C., Ambus, P., Milloux, M. J., & Andreux, F. (2007). Influence of <sup>15</sup>N enrichment on the net isotopic fractionation factor during the reduction of nitrate to nitrous oxide in soil. *Rapid Communications in Mass Spectrometry*, 21, 1447–1451.
- Milliken, G. A., & Johnson, D. E. (2009). Analysis of messy data. Volume 1. Designed experiments, New York: Dallas E. Chapman & Hall/CRC.
- Moore, K. J., & Dixon, P. M. (2015). Analysis of combined experiments revisited. Agronomy Journal, 107, 763–771.
- Nichols, V. A., Ordóñez, R. A., Wright, E. E., Castellano, M. J., Liebman, M., Hatfield, J. L., ... Archontoulis, S. V. (2019). Maize root distributions strongly associated with water tables in Iowa, USA. *Plant and Soil*, 444, 225–238.
- Ordóñez, R. A., Castellano, M. J., Hatfield, J. L., Helmers, J. M., Licht, M. A., Liebman, M., ... Archontoulis, S. V. (2018). Maize and soybean root front velocity and maximum depth in Iowa, USA. *Field Crops Research*, 215, 122–131.
- Pate, J. S., & Layzell, D. B. (1990). Energetics and biological costs of nitrogen assimilation. In B. J. Miflin & P. J. Lear (Eds.), *The biochemistry of plants* (Vol. 16, pp. 1–42). San Diego, CA: Academic Press.
- Pate, J. S., Unkovich, M. J., Armstrong, E. L., & Sanford, P. (1994). Selection of reference plants for <sup>15</sup>N natural abundance assessment of N<sub>2</sub> fixation by crop and pasture legumes in south-West Australia. *Australian Journal of Agricultural Research*, 45, 133–147.
- Peoples, M. B., Unkovich, M. J., & Herridge, D. F. (2009). Measuring symbiotic nitrogen fixation by legumes. In D. W. Emerich & H. B. Krishnan (Eds.), Nitrogen fixation in crop production (pp. 125–170). Madison, WI: ASA-CSSA-SSSA monograph.
- Rattalino Edreira, J. I., Mourtzinis, S., Conley, S. P., Roth, A., Ciampitti, I. A., Licht, M. A., Kandel, H., Kyveryga, P. M., Lindsey, L. E., Mueller, D. S., Naeve, S. L., Nafziger, E., Specht, J. E., Stanley, J., Staton, M. J., & Grassini, P. (2017). Assessing causes of yield gaps in agricultural areas with diversity in climate and soils. *Agricultural and Forest Meteorology*, 247, 170–180.
- Salvagiotti, F., Cassman, K. G., Specht, J. E., Walters, D. T., Weiss, A., & Dobermann, A. (2008). Nitrogen uptake, fixation and response to fertilizer N in soybeans: A review. *Field Crops Research*, 108, 1–13.
- Santachiara, G., Borrás, L., Salvagiotti, F., Gerde, J. A., & Rotundo, J. L. (2017). Relative importance of biological nitrogen fixation and mineral uptake in high yielding soybean cultivars. *Plant and Soil*, 418, 191–203.
- Setiyono, T. D., Cassman, K. G., Specht, J. E., Dobermann, A., Weiss, A., Yang, H., ... De Bruin, J. L. (2010b). Simulation of soybean growth and yield in near-optimal growth conditions. *Field Crops Research*, 119, 161–174.
- Setiyono, T. D., Walters, D., Cassman, K. G., Witt, C., & Dobermann, A. (2010a). Estimating maize nutrient uptake requirements. *Field Crops Research*, 118, 158–168.
- Setiyono, T. D., Weiss, A., Specht, J., Bastidas, A. M., Cassman, K. G., & Dobermann, A. (2007). Understanding and modeling the effect of temperature and daylength on soybean phenology under high-yield conditions. *Field Crops Research*, 100, 257–271.
- Shearer, G., & Kohl, D. H. (1986). N<sub>2</sub>-fixation in field settings: Estimations based on natural <sup>15</sup>N abundance. *Australian Journal of Plant Physiology*, 13, 699–756.
- Sinclair, T. R., & de Wit, C. T. (1975). Photosynthate and nitrogen requirements for seed production by various crops. *Science*, 189, 565–567.

- Sinclair, T. R., Farias, J. R., Neumaier, N., & Nepomuceno, A. L. (2003). Modeling nitrogen accumulation and use by soybean. *Field Crops Research*, 81, 149–158.
- Sinclair, T. R., & Rufty, T. W. (2012). Nitrogen and water resources commonly limit crop yield increases, not necessarily plant genetics. *Global Food Security*, 1, 94–98.
- Specht, J. E., Diers, B. W., Nelson, R. L., Toledo, F. J., Torrion, J. A., & Grassini, P. (2014). Soybean. In S. Smith, B. Diers, J. E. Specht, & B. F. Carver (Eds.), *Yield gains in major US field crops* (pp. 311–356). Madison, WI: ASA, CSSA, SSSA.
- Staswick, P. E. (1994). Storage proteins of vegetative plant tissues. Annual Review of Plant Biology and Plant Molecular Biology, 45, 303–322.
- Stephenson, R. A., & Wilson, G. L. (1977). Patterns of assimilate distribution in soybeans at maturity. I. The influence of reproductive development stage and leaf position. *Australian Journal of Agricultural Research*, 28, 203–209.
- Streeter, J., & Wong, P. P. (1988). Inhibition of legume nodule formation and N<sub>2</sub> fixation by nitrate. *Critical Reviews in Plant Sciences*, 7, 1–23.
- Tamagno, S., Balboa, G. R., Assefa, Y., Kovács, P., Casteel, S. N., Salvagiotti, F., ... Ciampitti, I. A. (2017). Nutrient partitioning and stoichiometry in soybean: A synthesis-analysis. *Field Crops Research*, 200, 18–27.
- Thibodeau, P. S., & Jaworski, E. G. (1975). Patterns of nitrogen utilization in the soybean. *Planta*, 127, 133–147.
- Thies, J. E., Singleton, P. W., & Bohlool, B. B. (1995). Phenology, growth, and yield of field-grown soybean and bush bean as a function of varying modes of N nutrition. *Soil Biology and Biochemistry*, 27, 575–583.
- Unkovich, M., Herridge, D., Peoples, M., Cadisch, G., Boddey, B., Giller, K., ... Chalk, P. (2008). *Measuring plant-associated nitrogen fixation in agricultural systems*. Canberra, Australia: Australian Centre for International Agricultural Research (ACIAR), Monograph number 136.
- Van Kessel, C., & Hartley, C. (2000). Agricultural management of grain legumes: Has it led to an increase in nitrogen fixation? *Field Crops Research*, 65, 165–181.
- Wang, E., & Engel, T. (1998). Simulation of phenological development of wheat crops. Agricultural Systems, 58, 1–24.
- Wilson, R. F. (2008). Soybean: Market driven research needs. In G. Stacey (Ed.), Genetics and genomics of soybean (pp. 3–15). New York, NY: Springer.
- Witty, J. F. (1983). Estimating N<sub>2</sub>-fixation in the field using <sup>15</sup>N-labelled fertilizer: Some problems and solutions. *Soil Biology and Biochemistry*, 15, 631–639.
- Yamagata, M., Kouchi, H., & Yoneyama, T. (1987). Partitioning and utilization of photosynthate produced at different growth-stages after anthesis in soybean (*Glycine max* L. Merr.): Analysis by long-term <sup>13</sup>Clabeling experiments. *Journal of Experimental Botany*, 38, 1247–1259.
- Yin, Y., Ying, H., Zheng, H., Zhang, Q., Xue, Y., & Cui, Z. (2019). Estimation of NPK requirements for rice production in diverse Chinese environments under optimal fertilization rates. *Agricultural and Forest Meteorology*, 279, 107756.

#### SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of this article.

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