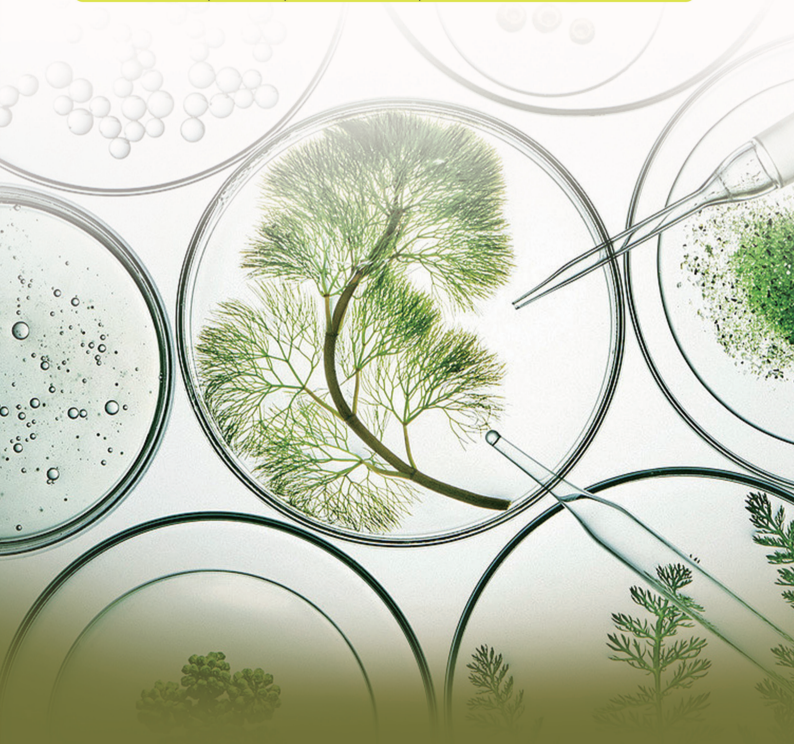


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ARTICLE

Diversity and Abundance of Amenity Trees in the Premises of International Institute of Tropical Agriculture (IITA), Ibadan, Nigeria

Yewande Owoeye^{1* (1)}, Stefan Hauser^{2 (1)}

ABSTRACT

Amenity trees are an essential element of most urban communities, contributing significantly to human well-being and improving environmental quality. Good knowledge of the diversity and abundance of trees in our environment and their importance can help promote conservation, which is essential for sustainability. This study aimed at assessing the diversity and abundance of amenity trees on the premises of the International Institute of Tropical Agriculture (IITA), Ibadan Nigeria. The institute was divided into working and residential areas. The trees in the study area were identified using a walking and windshield survey. A total population of 2626 trees from 126 species and 42 families were identified on the premises of IITA. The highest tree population of 523 trees was recorded in the Tropical Crescent residential area with 321 trees of *Lagerstroemia speciosa* being the most frequent species. Across working and residential areas, *Elaeis guineensis* was the most frequent species accounting for 19.92% of the total tree population. A Shannon-Wiener Diversity Index (H') of 3.383 and species evenness of 0.43 was obtained from the study area. The high values of diversity indices obtained indicate that IITA premises are rich in diverse tree species both indigenous and exotic hence should be referenced as a good urban landscape. The current management practices can be recommended for other institutions.

Keywords: Amenity trees; Species diversity; Abundance; IITA; Ibadan

1. Introduction

Amenity trees are trees that are not grown or

managed for their value as timber or crop but provide benefits or values [1]. Examples include trees found in parks and other recreational spaces, lining

*CORRESPONDING AUTHOR:

Yewande Owoeye, Forestry Research Institute of Nigeria, P.M.B, 5054, Ibadan, Nigeria; Email: yewandeowoeye@gmail.com

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¹ Forestry Research Institute of Nigeria, P.M.B, 5054, Ibadan, Nigeria

² International Institute of Tropical Agriculture, P.M.B, 5320, Ibadan, Nigeria

the sides of streets, railways, rivers, and canals and in home gardens. These trees have a positive impact on air quality through the deposition of pollutants in the vegetation, sequestration of atmospheric carbon dioxide in woody biomass, and reduction of temperature and associated ozone formation [2-4].

The use of trees in urban settings for different purposes cannot be overemphasized as they provide lots of benefits for humans and the environment at large. Their functions vary depending on the species, the site and the purpose for which they are planted. Trees are purposefully planted in academic environments/institutions for a variety of benefits, including aesthetics and other environmental services [5-8].

The presence of trees, particularly long-lived species that can withstand periodic reproductive constraints without obvious adverse demographic consequences, defines the landscape ^[9,10]. Over the years, over-exploitation has been a major environmental problem facing our forest reserves which has led to a drastic loss of tree diversity ^[11]. According to Omoro et al. ^[12], urbanization and infrastructure development have caused some disturbance to the trees that were purposefully planted in many urban communities, as well as in a variety of institutions such as hospital grounds, school or college campuses, and research institutes.

Knowledge of the composition, tree diversity, and species richness of tree populations in communities is crucial for the planning and implementation of biodiversity conservation efforts [9]. Also, understanding the diversity and distribution of trees in an urban setting would help provide information on the status of these trees. Over the years IITA, Ibadan campus has been reported to have a good and serene green environment comprising a large number of tree species, however, there are no scientific records on frequency, distribution and species composition in the working and residential area. Therefore, to promote tree conservation, especially in this institute, it is essential to evaluate the diversity and abundance of trees as this will guide the selection of tree species to be planted to increase benefits to the residents and contributes to the conservation of rare and threatened species.

2. Methodology

2.1 Study area

The International Institute of Tropical Agriculture was founded in 1967 in Ibadan, Nigeria. It is an award-winning research-for-development (R4D) organization providing solutions to hunger, poverty, and the degradation of natural resources in Africa. Prior to the acquisition of the 1000 ha of land by IITA through the Federal Government of Nigeria, the most extensive land use pattern was arable and tree crop farming and about 3000 people lived in about twenty-eight villages scattered in this area [13].

The IITA, Ibadan campus (**Figure 1**) has an area of 10 km², (1000 hectares) with about 92 hectares of residential and working area (**Figure 2**). It is located between latitude 7°29'16.76" and 7°30'12" N; and between longitude 3°54'50.30" and 3°52'43.55" E and at an altitude of 227 m above sea level.

The rainfall pattern is bimodal with an annual total ranging from 1,300-1,500 mm most of which falls between May and September. The average daily temperature ranges between 21 °C and 23 °C while the maximum is between 28 °C and 34 °C. Radiation is about 5285 MJ/m²/year. The mean relative humidity is in the range of 64% to 83% [14].

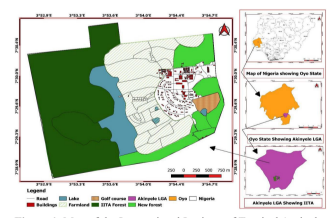


Figure 1. Map of the International Institute of Tropical Agriculture.

Source: Produced from QGIS, 2021.

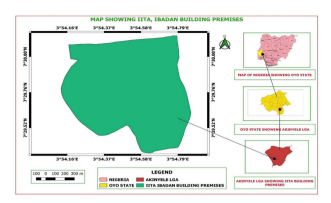


Figure 2. Map of the residential and working areas of the International Institute of Tropical Agriculture.

Source: Produced from QGIS, 2021.

2.2 Method of data collection

This study covers the working and residential premises of the International Institute of Tropical Agriculture, (IITA), Ibadan campus (**Figure 2**). Trees within these premises were counted and identified to species level using walking and wind shield survey methods. The structure of the amenity tree population such as species composition, frequency, and relative frequency was recorded.

2.3 Data analysis

Relative Density (RD) was obtained using the formula given by Oduwaiye et al. [15]:

$$RD = \frac{n_i}{N} \times 100$$

where RD = relative density, n_i = number of individuals of species and N = total number of individuals in the entire population.

Tree population and diversity were analyzed using the following species diversity indices.

A mathematical technique that takes into account the species richness and abundance was used to determine tree diversity. The Shannon-Wiener diversity index was calculated using the following equation provided by Price [16]:

$$H^1 = \sum s i=1 \ pi \ Lnpi....$$

H¹ is the Shannon diversity index, S is the total number of species in the sample plot, pi is the proportion of a species relative to the total number of plants in the sample plots and Ln is the natural logarithm.

Species evenness (E) in each of the plots was determined using Shannon's equitability (E_H) as stated by Kent and Coker [17]:

$$E = \frac{H^1}{Ln(S)}$$

S is the total number of species in each premises (working and residential premises).

3. Results

Tree species diversity and abundance in the study area are presented in **Table 1**. The working and residential areas contained a total of 2626 individual trees from 126 different tree species from 42 different families. The Shannon-Wiener index of diversity was 3.38 and the species evenness was 0.43.

Table 2 shows the origin, frequency and relative frequency of tree species encountered in the study area. The oil palm *Elaeis guineensis, Lagerstroemia speciosa, Roystonea regia and Tabebuia rosea* were present at frequencies (and relative frequencies) of 523 (19.92%), 321 (12.22%), 217 (8.26%) and 173 (6.59%), respectively. Of all the species encountered, *Elaeis guineensis* was the most frequent in the study area.

Table 3 shows the family distribution of the species encountered on the premises of IITA, Ibadan. *Arecaceae* were the most frequent species with a relative frequency of 32.83% followed by *Fabaceae* with 13.37% and *Lythraceae* with 12.22%.

Table 1. Tree species diversity and abundance in the study area.

Diversity indices	Values		
Total No. of individual	2626		
No. of individual species	126		
No. of family	42		
Shannon diversity index (H1)	3.383		
Evenness	0.43		

Table 2. Frequency, origin, family and relative frequency of individual trees by species on IITA, Ibadan campus.

S/N	Tree species	Origin	Family	Frequency	Relative frequency (%)	
1	Adansonia digitata	I	Malvaceae	1	0.04	
2	Afzelia africana	I	Fabaceae	1	0.04	
3	Albizia ferruginea	I	Fabaceae	2	0.08	
4	Albizia glaberrima	I	Fabaceae	2	0.08	
5	Albizia lebbeck	Е	Fabaceae	2	0.08	
6	Albizia saman	Е	Fabaceae	1	0.04	
7	Albizia zygia	I	Fabaceae	13	0.50	
8	Allophylus africanus	I	Sapindaceae	2	0.08	
9	Alstonia boonei	I	Apocynaceae	3	0.11	
10	Anacardium occidentalis	Е	Anacardiaceae	1	0.04	
11	Anogeisus leiocarpus	I	Combretaceae	2	0.08	
12	Anthocleista djalonensis	I	Loganiaceae	1	0.04	
13	Anthocleista nobilis	I	Gentianaceae	6	0.23	
14	Anthocleista vogelii	I	Loganiaceae	2	0.08	
15	Anthonotha macrophylla	I	Caesalpinioideae	2	0.08	
16	Antiaris toxicaria	I	Moraceae	6	0.23	
17	Araucaria heterophylla	Е	Araucariaceae	3	0.11	
18	Artocarpus altilis	Е	Moraceae	8	0.30	
19	Artocarpus communis	Е	Moraceae	2	0.08	
20	Artocarpus heterophyllus	Е	Moraceae	4	0.15	
21	Asimina tribola	Е	Annonaceae	4	0.15	
22	Azadirachta indica	Е	Meliaceae	7	0.27	
23	Bauhinia purpurea	Е	Fabaceae	5	0.19	
24	Bauhinia variegata	Е	Fabaceae	2	0.08	
25	Blighia sapida	I	Sapindaceae	15	0.57	
26	Bombacopsis glabra	Е	Bombacaceae	1	0.04	
27	Bombax buonopozense	I	Malvaceae	2	0.08	
28	Borassus aethiopum	I	Arecaceae	14	0.53	
29	Bosqueia angolensis	I	Moraceae	4	0.15	
30	Brachystegia eurycoma	I	Fabaceae	8	0.30	
31	Calliandra haematocephala	Е	Fabaceae	5	0.19	
32	Callophyllum macrocarpum	Е	Clusiaceae	4	0.15	
33	Carica papaya	Е	Caricaceae	5	0.19	
34	Cassia fistula	Е	Fabaceae	8	0.30	
35	Cassia javanica	Е	Fabaceae	6	0.23	
36	Cassia nodosa	Е	Fabaceae	3	0.11	
37	Casuarina equisetifolia	Е	Casuarinaceae	11	0.42	
38	Ceiba pentandra	I	Malvaceae	8	0.30	
39	Chrysophyllum albidum	I	Sapotaceae	4	0.15	

Table 2 continued

S/N	Tree species	Origin	Family	Frequency	Relative frequency (%)
40	Citrus reticulata	Е	Rutaceae	33	1.26
41	Citrus sinensis	Е	Rutaceae	3	0.11
42	Cleistopholis patens	I	Annonaceae	4	0.15
43	Cnetis ferruginea	I	Connaraceae	2	0.08
44	Cocos nucifera	Е	Arecaceae	31	1.18
45	Cola acuminata	I	Malvaceae	2	0.08
46	Cola gigantea	I	Malvaceae	3	0.11
47	Cola nitida	I	Malvaceae	40	1.52
48	Dacryodes edulis	I	Burseraceae	6	0.23
49	Dactyladenia barteri	I	Chrysobalanceae	1	0.04
50	Daniellia oliveri	I	Fabaceae	6	0.23
51	Delonix regia	Е	Fabaceae	36	1.37
52	Dichrostachys cinerea	I	Fabaceae	6	0.23
53	Dobera glabra	I	Salvadoraceae	10	0.38
54	Duranta repens	Е	Verbenaceae	3	0.11
55	Elaeis guineensis	I	Arecaceae	523	19.92
56	Entandrophragma angolense	I	Meliaceae	1	0.04
57	Enterolobium cyclocarpum	Е	Fabaceae	7	0.27
58	Erythrina variegata	Е	Fabaceae	1	0.04
59	Erythrophleum suaveolens	I	Leguminosae	1	0.04
60	Eucalyptus camaldulensis	Е	Myrtaceae	4	0.15
61	Eucalyptus tereticornis	Е	Myrtaceae	33	1.26
62	Eugenia uniflora	Е	Myrtaceae	3	0.11
63	Ficus aurea	Е	Moraceae	2	0.08
64	Ficus benjamina	Е	Moraceae	3	0.11
65	Ficus exasperata	I	Moraceae	4	0.15
66	Ficus lutea	I	Moraceae	27	1.03
67	Ficus mucuso	Е	Moraceae	1	0.04
68	Gliricidia sepium	Е	Fabaceae	7	0.27
69	Gmelina arborea	Е	Lamiaceae	1	0.04
70	Hildegardia barteri	I	Malvaceae	44	1.68
71	Holarrhena floribunda	I	Apocynaceae	5	0.19
72	Hura crepitans	Е	Euphorbiaceae	34	1.29
73	Irvingia gabonensis	I	Irvingiaceae	2	0.08
74	Jacaranda mimosifolia	Е	Bignoniaceae	3	0.11
75	Kigelia africana	I	Bignoniaceae	1	0.04
76	Lagerstroemia speciosa	Е	Lythraceae	321	12.22
77	Leucaena leucocephala	Е	Fabaceae	9	0.34
78	Mangifera indica	Е	Anacardiaceae	28	1.07
79	Milicia excelsa	I	Moraceae	12	0.46

Table 2 continued

S/N	Tree species	Origin	Family	Frequency	Relative frequency (%)
80	Milletia thonningii	I	Fabaceae	10	0.38
81	Monodora myristica	I	Annonaceae	6	0.23
82	Monodora teluifolia	I	Annonaceae	24	0.91
83	Morinda lucida	I	Rubiaceae	2	0.08
84	Moringa oleifera	I	Moringaceae	2	0.08
85	Myrianthus arboreus	I	Urticaceae	1	0.04
86	Nauclea diderrichii	I	Rubiaceae	1	0.04
87	Newbouldia laevis	I	Bignoniaceae	6	0.23
88	Parkia biglobosa	I	Fabaceae	2	0.08
89	Peltophorum pterocarpum	Е	Fabaceae	72	2.74
90	Persea americana	Е	Lauraceae	9	0.34
91	Phoenix reclinata	I	Arecaceae	77	2.93
92	Pinus caribaea	Е	Pinaceae	50	1.90
93	Plumeria alba	Е	Apocynaceae	7	0.27
94	Plumeria rubra	Е	Apocynaceae	14	0.53
95	Polyalthia longifolia	Е	Annonaceae	90	3.43
96	Psidium guajava	Е	Myrtaceae	6	0.23
97	Pterocarpus osun	I	Leguminosae	4	0.15
98	Pterocarpus santalinoides	I	Fabaceae	3	0.11
99	Pterocarpus soyauxii	I	Fabaceae	6	0.23
100	Pterospermum heterophyllum	Е	Malvaceae	6	0.23
101	Pycnanthus angolensis	I	Myristicaceae	7	0.27
102	Pyrus communis	Е	Rosaceae	1	0.04
103	Rauvolfia vomitoria	I	Apocynaceae	2	0.08
104	Ravenala madagascariensis	I	Strelitziaceae	6	0.23
105	Ricinodendron heudolotii	I	Euphorbiaceae	1	0.04
106	Roystonea regia	Е	Arecaceae	217	8.26
107	Senna alata	Е	Fabaceae	1	0.04
108	Senna fistula	Е	Fabaceae	127	4.84
109	Spathodea campanulata	I	Bignoniaceae	6	0.23
110	Sterculia tragacantha	I	Sterculiaceae	6	0.23
111	Syzygium malaccense	Е	Myrtaceae	115	4.38
112	Tabebuia rosea	Е	Bignoniaceae	173	6.59
113	Tectona grandis	Е	Lamiaceae	9	0.34
114	Terminalia catappa	I	Combretaceae	43	1.64
115	Terminalia ivorensis	I	Combretaceae	1	0.04
116	Terminalia mantaly	Е	Combretaceae	4	0.15
117	Terminalia superba	I	Combretaceae	5	0.19
118	Theobroma cacao	Е	Malvaceae	12	0.46
119	Treculia africana	I	Moraceae	6	0.23

Table 2 continued

S/N	Tree species	Origin	Family	Frequency	Relative frequency (%)
120	Trema orientalis	I	Cannabaceae	17	0.65
121	Trichilia africana I		Meliaceae	5	0.19
122	Trichilia megalantha	I	Meliaceae	2	0.08
123	Trichilia monadelpha	I	Meliaceae	12	0.46
124	Trilepisium madagascariense	I	Moraceae	3	0.11
125	Triplochiton scleroxylon	I	Malvaceae	3	0.11
126	Zanthoxylum zanthoxyloides	I	Rutaceae	1	0.04
	Total			2626	100.00

Source: Field survey 2021.

 Table 3. Family distribution of tree species in the study area.

S/N	Family	No. of SPP/Family and abundance	Relative frequency
1	Anacardiaceae	29	1.10
2	Annonaceae	128	4.87
3	Apocynaceae	31	1.18
4	Araucariaceae	3	0.11
5	Arecaceae	862	32.83
6	Bignoniaceae	189	7.20
7	Bombacaceae	1	0.04
8	Burseraceae	6	0.23
9	Caesalpinioideae	2	0.08
10	Cannabaceae	17	0.65
11	Caricaceae	5	0.19
12	Casuarinaceae	11	0.42
13	Chrysobalanceae	1	0.04
14	Clusiaceae	4	0.15
15	Combretaceae	55	2.09
16	Connaraceae	2	0.08
17	Euphorbiaceae	35	1.33
18	Fabaceae	351	13.37
19	Gentianaceae	6	0.23
20	Irvingiaceae	2	0.08
21	Lamiaceae	10	0.38
22	Lauraceae	9	0.34
23	Leguminosae	5	0.19
24	Loganiaceae	3	0.11
25	Lythraceae	321	12.22
26	Malvaceae	121	4.61
27	Meliaceae	27	1.03
28	Moraceae	82	3.12
29	Moringaceae	2	0.08

Table 3 continued

S/N	Family	No. of SPP/Family and abundance	Relative frequency
30	Myristicaceae	7	0.27
31	Myrtaceae	161	6.13
32	Pinaceae	50	1.90
33	Rosaceae	1	0.04
34	Rubiaceae	3	0.11
35	Rutaceae	37	1.41
36	Salvadoraceae	10	0.38
37	Sapindaceae	17	0.65
38	Sapotaceae	4	0.15
39	Sterculiaceae	6	0.23
40	Strelitziaceae	6	0.23
41	Urticaceae	1	0.04
42	Verbenaceae	3	0.11
Total		2626	100.00

4. Discussion

The IITA Ibadan premises comprise a diverse set of indigenous and exotic tree species. The surveyed area of 92 ha containing 2626 trees translates to a density of 28.54 trees per ha. Unfortunately, there is no data on tree densities for other African Institutions or parks to compare this result. The trees in IITA are generally in good condition with a few exceptions of exotic species. The placement of the trees serves largely the purpose of shading in the residential area with a good portion being used as fruit trees. The placement in the working area is more for ornamental purposes and to line streets and parking areas. Many trees are old and have been exposed to recent bad weather events. IITA does replace dead and damaged trees and does conduct management operations to retain trees in good condition. IITA staff and residents consider the presence of the trees as beneficial irrespective of the trees' purpose and use. Most of the trees are beneficial to the people on the premises as they provide shade and fruit and improve the aesthetics, which are some of the important benefits of amenity trees. The total of 2626 individual trees; with 126 different tree species from 42 different families compares positively with the results obtained by Agbelade et al. [18] on the 'Assessment of Urban Forest Tree Species Population and Diversity in Ibadan, Nigeria', who recorded a total number of 155 individual trees belonging to 16 families. This implies that the study area, despite being a small fragment of Ibadan city is well diversified comprising more tree species at a higher population density than other similar sites. The abundance and diversity of trees in the IITA area can be linked to the policy of conservation guiding the management of trees on the premises. IITA has a nursery of indigenous African trees and focuses on propagating rare and threatened species to contribute to the conservation of such species.

The benefits of the trees to the staff and residents on the premises are variable and largely to provide shade (e.g. Lagerstroemia speciosa, Roystonea regia, Senna fistula, Peltophorum pterocarpum), provision of food/fruits (e.g. Citrus spp, Elaeis guineensis, Dacryodes edulis, Cocos nucifera, Syzygium malaccense, Persea americana) improving the aesthetics (e.g. Polyalthia longifolia, Roystonea regia, Ravenala madagascariensis).

The study area is dominated by the family Arecacea with 862 individuals, mostly oil palm (*Elaeis guineensis*) which are almost all remnants of the previous land use by the smallholders using the land before the installation of IITA. These palms are still harvested and maintained by the successors of the villagers who were displaced. The still high numbers

of oil palm are thus due to the continued use that has prevented any felling by the current residents. Other dominant families include Fabaceae with 351 individuals, mostly serving aesthetic purposes and Lythraceae having 321 individuals.

The Shannon Diversity Index (H') in the study area was 3.38, which indicates a high tree species diversity. These findings are similar to the results obtained by Haastrup et al. [19] for the Owo forest reserve and Ogundele et al. [20] for the Akure forest reserve with species diversity of 3.42 and 3.18, respectively. The H' value of this study is similar to the one recorded by Agbelade et al. [18] who recorded a 3.35 Shannon Diversity Index (H'). Across the available data, it can be postulated that the tree species diversity in the work and residential areas of IITA is amongst the highest recorded in southwest Nigeria.

5. Conclusions

The results obtained from this study revealed the diversity and abundance of trees in the working and residential areas of the IITA, Ibadan campus. The study area comprised a lot of tree species when compared with the results of studies obtained in the Ibadan metropolis and some other forest reserves in southwest Nigeria. The species in this premise are said to have been well conserved and managed over the years. Some of the indigenous trees which are now endangered were encountered on this site, therefore, this study area can be said to be a home of different species which are of great importance. The various species encountered during this study revealed that IITA, Ibadan is a good example of urban green space; hence, the results from this study can be used in developing the database required for urban green space management.

Authors' Contributions

Dr. Stefan Hauser facilitated access to the premises, liaised with IITA management and residents to obtain permission to enter the working area and private residential spaces, reviewed the manuscript.

Yewande Owoeye collected and compiled the

data on amenity trees from the International Institute of Tropical Agriculture IITA, Ibadan.

Conflict of Interest

The authors declare no conflict of interest.

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ARTICLE

Effect of Alternate Bearing Phenomenon and Boron Foliar Application on Nitrogen-15 Uptake, Translocation and Distribution in Mango Tree (cv. Zebda)

Rawia El-Motaium^{1*}, Ayman Shaban², El Sayed Badawy³, Ahmad Ibrahim¹

ABSTRACT

The objectives of this investigation are to study nitrogen uptake, translocation, accumulation and distribution in mango tree organs using labeled nitrogen (15 N) and to understand the mechanism of boron action in increasing fruit yield in the off-year. A field experiment was conducted using fifteen-year-old mango trees (cv. Zebda) grown at Al Malak Valley Farm, El-Sharkeya Governorate-Egypt. Treatments included the application of (15 NH₄) $_2$ SO₄, "in the on-year", at a rate of 50 g nitrogen/ tree through the stem injection technique. While boron was sprayed on the same trees "in the off-year" at the following rates: 0.0 (control), 250 and 500 mg·L⁻¹. The authors hypothesize that boron and nitrogen act synergistically to increase mango fruit yield in the off-year. Results indicated that the highest 15 N uptake and accumulation in the on and off-years was observed in the upper (young leaves). When boron was applied at 250 mg·L⁻¹, in the off-year, the upper (young leaves) recorded the highest 15 N uptake and accumulation (6 15Ndff = 13.93) relative to the other two leaf categories and those of the on-year. In the on-year fruit accumulated higher 15 N than leaf or bud. In the off-year, bud exhibited the highest 15 N accumulation without boron application, while leaves exhibited the highest 15 N with boron application. The highest 6 15Ndff in all tree organs was observed at 250 mg·L⁻¹ boron rate. Boron increased nitrogen uptake, translocation and accumulation in mango tree organs. A synergistic relationship was observed between boron and nitrogen which led to an increase in fruit yield in the off-year.

Keywords: Mango; ¹⁵N distribution; ¹⁵N-stem injection technique; ¹⁵N translocation; ¹⁵N uptake, ¹⁵N accumulation; On and off-year; Synergistic relationship

*CORRESPONDING AUTHOR:

Rawia El-Motaium, Plant Researches Department, Nuclear Research Center, Atomic Energy Authority, Inshas, Cairo, 13759, Egypt; Email: el-motaiumr@yahoo.com

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¹ Plant Researches Department, Nuclear Research Center, Atomic Energy Authority, Inshas, Cairo, 13759, Egypt

²Department of Pomology, Faculty of Agriculture, Cairo University, Giza, Cairo, 12613, Egypt

³ Soil Science Department, Faculty of Agriculture, Cairo University, Giza, Cairo, 12613, Egypt

1. Introduction

Mango (*Mangifera indica* L.) is one of the most popular fruit crops in Egypt and is of great economic importance. Irregularity of flowering in mango, which varies in time and intensity from year to year, is a common phenomenon [1]. Mango has many problems related to fruit set, yield and quality due to the imbalance of nutrient supply [2]. Mango is an extremely alternate bearing species and varieties vary in their alternation degree. The "Zebda" is classified as being a highly alternate bearing cultivar with low productivity due to this phenomenon. Strategies to enhance yield in the off-year or to develop more consistent fruiting in the on- and off-years would improve grower returns.

Alternate-bearing trees are those trees that do not bear a regular crop year after year, but rather a heavy yield followed by an extremely light one and vice versa. There are several theories of the alternate bearing phenomenon. The alternation phenomenon is caused by environmental triggers and endogenous factors, which lead to a shift in the balance between vegetative and reproductive growth [3]. The C/N ratio was reported to influence fruit set and consequently yield, rather than flowering [4-6]. Regulatory roles in various phases of the alternate bearing cycle have been related to plant hormones [7], mineral nutrients [8] and carbohydrates [9,10].

Boron has a crucial function in improving flower fertility and fruit set ^[11]. This is mainly due to the role of boron in increasing pollen grains germination and pollen tube elongation which leads to higher fruit set and yield ^[12,13] and/or to an increase in the flavonoid content of the pollen ^[14]. Boron increases the initial and final fruit set in almonds ^[15]. Boron deficiency can result in a reduction of fruit quality and yield potential in mangoes ^[16]. The increase in fruit set and yield in mango due to boron foliar application was emphasized by several researchers ^[17-24]. Thus, boron may help to solve the alternate bearing phenomena by promoting flowering and fruit set in the off-year.

Application of nitrogen fertilizer at the proper rate and time may reduce irregular bearing in mangoes. Significant increase in blooming occurred in the off-year and flushing in the on-year when mango trees received 3% urea ^[25]. Nitrogen increases leaf chlorophyll levels and photosynthesis, thus promoting shoot growth and flowering. Spraying "Langra" mango trees with urea at 2% decreased flowering percentage in the "on-year" considerably ^[26]. Sprayed 15-year-old mango trees with urea solution (0.8%) increased panicle length and number of male and perfect flowers ^[27].

Nitrogen and boron when applied together may be involved in the alleviation of alternate bearing. Boron in combination with urea significantly increased the flowering and fruiting of mango "Fazli" particularly at a 0.4% rate [28]. Along the same line, several researchers reported a positive interaction "synergistic" between boron and nitrogen. It was reported by Miley et al. [29] that boron enhanced the utilization of nitrogen by increasing the translocation of nitrogen compounds into the cotton boll. Davis et al. [30] found that in field culture, boron application (foliar and/or soil) resulted in an increase in boron and nitrogen uptake and tissue concentration by tomato plants.

The synergistic effects of B and N may be a consequence of enhanced nitrate assimilation and nitrate reductase activity as shown in Tobacco plants grown with differential boron application [31]. The effect of nitrogen and boron fertilizer on the alternate bearing cycle in Pistachio trees was investigated by Weinbaum et al. [32]. Their results showed that the on-year trees have greater reproductive demand for N and carbohydrates, reduced accumulation of C and N reserves and reduced recovery of applied labeled-N fertilizer than the off-year trees.

The allocation pattern of ¹⁵N in leaves of different ages was studied by Biddulph ^[35] who stated that the different ¹⁵N distributions found in leaves of different ages are probably due to the fact that the leaves differ in their basic metabolism according to age. In young leaves, the predominant process is the synthesis of new protoplasm, which results in growth towards maturity. In older, mature leaves, photosynthesis is the dominant function, and little growth

takes place.

Burr and Takahashi [36] demonstrated the uneven distribution of the atomic percentage of ¹⁵N in the tissue of different origins and ages with the highest ¹⁵N accumulation in leaves that were at the stage of maximum activity. They added that nitrogen flows to places where there is metabolic demand and not to those where there is a nutritional vacuum. It was suggested that the redistribution of nitrogen in the plant is the result of competition between meristems and other tissues of different metabolic activity ^[37].

The distribution of ¹⁵N in plants might give valuable information about the extent and rate of dynamic exchange (breakdown vs. resynthesis) of cellular proteins in tissues of various ages, as well as from different plant organs ^[38]. The distribution of ¹⁵N-labeled fertilizer applied to Pecan trees was studied by Kraimer et al. ^[39]. They found that early spring growth, flowering and embryo development used fertilizer-N applied the previous year as well as that applied during the current year.

El-Motaium et al. [40] found that using the stem injection technique in olive trees, ¹⁵N a.e. in leaves and flowers appear to increase in response to boron foliar application. The distribution of ¹⁵N between the leaves and flowers of the olive tree indicated a higher enrichment of ¹⁵N in the flowers than in the leaves. The maximum increase in %Ndff was observed at 200 mg·L⁻¹ boron application rate for both leaves and flowers. This boron dose (200 mg·L⁻¹) coincides with the highest values for flowering; fruit set and yield [41]. They suggested that boron is likely required for the synthesis of certain nitrogen compounds. This might be nucleic acid particularly the nitrogen base "uracil" [42] that enhances flowering and fruit set processes. Boron has been shown to increase flowering and fruiting by increasing pollen grains germination and pollen tube elongation [43,44,12] which can lead to higher fruit set and yield.

Several methods have been tested for using the ¹⁵N tracer technique with trees including soil and foliar application ^[33]. Soil fertilization often results in soil contamination with ¹⁵N and the residual ¹⁵N complicates the estimation of reserve use in the sec-

ond season. Foliar fertilization is neither adequate nor uniform to large trees which requires a large amount of solution. The stem injection technique was proposed to be the most suitable method for labeling trees as it is a non-destructive technique. Injection fertilization of fruit trees has several advantages over soil fertilization [34]. These advantages are: There is no fertilizer loss to the groundwater, only 5-10% of the label N applied to the soil is needed, and weed control is not needed since weed roots do not compete with tree roots for nutrient uptake. In addition, it labels the tree without affecting the soil N pools.

Although biennial bearing has been reported to be genetically controlled, the physiological factors governing such bearing habit have not been clearly understood [45,46]. Therefore, the objective of this research is to study the effect of the on and off years and boron foliar spray on nitrogen uptake, translocation and accumulation in mango trees in an attempt to shed light on the physiological and biochemical basis of boron's role in increasing flowering, fruit set and yield in the off-year.

2. Materials and methods

Fifteen years old mango (*Mangiferaindica*) trees (cv. Zebda) grown at Al Malak Valley Farm, El-Sharkeya Governorate-Egypt (30-51° North; 32-53° East) were used in this field experiment. The trees were grown in sandy soil, "Typic Torripsamments" [47] with total N = 0.06%, total $B = 19.4 \text{ mg} \cdot \text{kg}^{-1}$, available $B = 0.51 \text{ mg} \cdot \text{kg}^{-1}$) under drip irrigation system.

2.1 Preparation of labeled nitrogen fertilizer (N-15)

Labeled ammonium sulfate fertilizer was applied to mango trees at 50 g N/tree rate once in January of the on-year. The fertilizer was dissolved in 500 mL of deionized water. The labeled fertilizer, (¹⁵NH₄)-₂SO₄, is enriched with 10.35% ¹⁵N atom excess. The (¹⁵NH₄)₂SO₄ solution was loaded into the trees' xylem vessels through the trunk using the stem injection technique.

2.2 Stem injection technique

The injection of (¹⁵NH₄)₂SO₄ solution was conducted according to the following steps and as shown in **Figure 1**:

- A circular disk of about 2-3 cm diameter was removed from the bark at the base of the trunk;
- In the middle of the removed disk a pore of about 1 cm in diameter was made at the base of the trunk, 15 cm from the ground at a 45° angle, and through about 75% of the tree diameter;
- A hard plastic tube was inserted in the pore and tightened with plastic material;
- An injection needle was tightly connected to a 500 mL tank (reservoir) containing the N-15 fertilizer solution (50 g 15 N) + CuSO₄(0.588 g) to avoid introducing pathogens into the tree;
- The tank was located 1 m higher than the injection hole.

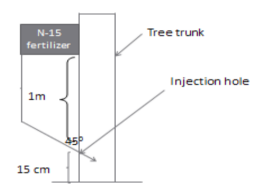


Figure 1. Diagram of N-15 stem injection technique.

The only source of N-15 in this experiment is the injection solution in the on-year. While, in the off-year, the reserved and recycled ¹⁵N within the tree was the source of the labeled nitrogen. Our results show that the N-15 enriched [(¹⁵NH₄)₂SO₄] at 50 g N/ tree rate was sufficient to label the mango trees and to detect the ¹⁵N in all the tree organs in both the on and off years.

Boron was sprayed as boric acid (H₃BO₃) in the same trees in January of the following year (off-year), using tractor mounted sprayer at the following rates: 0.0 (control), 250 and 500 mg·L⁻¹. Tween 20 was added to the foliar solution at a 1 mL/L rate. Each tree was sprayed with 15 litres of the boron solution

to achieve runoff.

2.3 Leaves and fruit sampling

Leaf and bud samples of the on-year were collected in March, two months from the initiation of the ¹⁵N injection following the depletion of the injection solution. At the commencement of the off-year, leaves were selected from the same relative position to leaves of the on-year. Leaf and bud samples were collected in February, one month after the boron spray. Leaves were collected in three categories according to their position on the branch and age, upper leaf (3-month-old), middle leaf (6-month-old) and lower leaf (10-month-old). In the on-year and off-year fruits were collected in August. All samples were dried at 70 °C then ground up to very fine powder.

Leaf, bud and fruit were analyzed for %¹⁵N atom excess and total nitrogen content. The percentage of ¹⁵N atom excess was determined using the Emission Spectrometry N-15 Analyzer (FAN Fisher No. 1-6PC Spectrometer). Total nitrogen was determined using the C/N Analyzer (Elementar, Germany).

2.4 Calculation

The following equation was used to calculate nitrogen percentage in the plant organs derived from fertilizer (%Ndff) according to Zapata [48]:

%N dff = (% 15N a. e. in plant sample / % 15N a. e. in the labeled fertilizer) x100

2.5 Soil analysis

Soil physical and chemical properties

Soil samples were collected from the soil profile at (0-25 and 25-50 cm) depth to determine the following characteristics, results are shown in **Table 1**:

- pH and EC using pH meter and conductivity meter, respectively (Cole Parmer, USA);
 - Calcium carbonate (CaCO₃) using Calcimeter;
 - Bulk density (BD) according to Mckenzie et al. [49];
- Organic matter (OM) using the Walkley-Black procedure, according to Chapman and Pratt [50];

- Texture class, according to Dewis and Freitas [51]. *Soil nutrients analysis*

Soil available boron was extracted by the hot water extract method according to Johnson and Fixen ^[52]. Soil total boron was extracted by (5 mL HNO₃ + 2 mL HF + 2 mL HCL acids) using the Microwave Digestion System (Milestone, Italy). Boron was determined by the Azomethine dye method according to Johnson and Fixen ^[52]. Soil total nitrogen was determined using the C/N Analyzer (Elementar, Germany). Soil phosphorus was determined using the vanadate-molybdate method ^[53] and measured using the UV-VIS Spectrophotometer (Shimadzu, Japan) at 430 nm. Soil potassium was determined using the Flame-Atomic Absorption Spectrometry (Shimadzu 6800, Japan).

2.6 Flowering, fruit set and yield analysis

Flower percentage

Counting the number of flowers in a mango tree is complex, because of the huge number of flowers in addition to the difficulty of distinguishing between male flowers and Hermaphrodite flowers. Therefore, an indication of flowering percentage was used. Twenty-one-year-old branches were chosen for every

tree. Flower percentage was calculated at the end of April using the following equation:

Flower percentage = (the number of flowering branches / the total number of labeled brances) x 100

Final fruit set percentage

The final fruit set was determined as the number of retained fruits per panicle at harvest relative to the initial fruit set according to Shaban [54] and calculated using the following equation:

Final fruit set = [(the number of retained fruit per panicle at harvest / the number of initial fruit set)] x 100

Yield

Fruits were collected at the maturity stage. Each individual tree was harvested manually and fruit weight was estimated. The average weight for each treatment was calculated. Yield was expressed as kg fruit/tree.

2.7 Experimental design and statistical analysis

Treatments were arranged in a completely randomized block design with three replicates for each treatment and nine trees per block. Statistical analysis was performed using the MSTAT microcomputer program. The significant means were compared using LSD at 5% probability according to Snedecor and Cochran [55].

Boron Coarse Fine Organic Silt Clav Texture EC TCC BD N (mg·kg⁻¹) Depth sand sand pН matter (dS/m) class (g/cm^3) (%)(%) (%)(%) (%) (%) (%)Total Available 0-25 67.9 28.9 1.25 7.56 0.34 1.55 1.64 0.07 19.7 0.53 1.88 Sandy 13.0 25-50 63.5 33.1 2.05 1.36 Sandy 7.81 0.30 1.00 15.0 0.05 19.1 0.50

Table 1. Soil physical, chemical and nutrient characteristics.

3. Results and discussion

The fate of the labeled nitrogen (¹⁵N) in the mango tree organs is shown in **Tables 2-5**. Results are expressed as an average of two successive seasons.

3.1 ¹⁵N uptake, translocation, distribution and total nitrogen percentage in mango tree organs during the "on-year"

Results in Table 2 show that the %Ndff values

are 7.41 for the upper leaves, 6.10 for the middle leaves and 5.23 for the lower leaves. The buds showed a %Ndff value of 6.28 while fruits had the highest translocated ¹⁵N enrichment of 12.08% and 8.99% for peel and pulp, respectively. The ¹⁵N distribution pattern was highest in fruit but lowest in leaf.

Results in **Table 2** show that the descending order of %¹⁵Ndff is as follows: Fruit is 12.08 peel and 8.99 pulp, upper leaves is 7.41, buds is 6.28, medium leaves is 6.10 and lower leaves is 5.23. Total

nitrogen percent was highest in the old lower leaves (1.24%) then it decreased with decreasing the leaf age to reach 1.19% for the middle leaves and 1.05% for the upper leaves. The labeled N in the fruit of the on-year represents 84% of the total labeled N accumulated in leaf + bud.

The current study shows that in the on-year, the difference in %¹⁵Ndff between the different leaves categories (young, medium, old) is due to their position and age. The high %¹⁵Ndff shown by the upper young leaves could be due to they were at their maximum activity stage. The on-year trees showed higher ¹⁵N enrichment in the reproductive organ (fruit) than in the vegetative organ (leaf). This finding supports the idea of Burr and Takahashi ^[36] who stated that nitrogen flows to places where there is high metabolic demand (activity). Fruits in the on-year represent the organ of high metabolic demand (preferential sink).

In the on-year, the distribution pattern of the labeled (¹⁵N), was not uniform among mango tree organs. Higher ¹⁵N enrichment was observed in young leaves than the buds which indicates that young leaves have higher metabolic demand for nitrogen than buds. Fruit was the dominant sink for nitrogen. The results agree with Weinbaum et al. ^[32]. The uneven distribution of ¹⁵N could be due to the slow translocation and redistribution of mobile nitrogen in the tree ^[38]. It also indicates that the stationary state of the breakdown-resynthesis turnover of nitrogen is still not reached ^[38]. Our results agree with Wallace et al., ^[56] who found that the uneven distribution of ¹⁵N was observed in plant parts after 60 days of ¹⁵N-labeled (NH₄)₂SO₄ application.

3.2 Influence of boron on ¹⁵N accumulation in mango tree organs during the "off-year"

In the absence of boron application (zero boron), the greatest 15 N translocation occurred to the upper leaves ($\%^{15}$ Ndff = 5.24) compared with the middle (($\%^{15}$ Ndff = 4.34) or the lower (($\%^{15}$ Ndff = 3.86) leaves. With boron application at (250 mg·L⁻¹) the $\%^{15}$ Ndff increased by almost three fold in all leaf categories compared with the control (**Table 3**). The up-

per leaf (13.93) still maintained the highest %¹⁵Ndff values compared with the middle (12.95) and lower leaf (10.86). Boron treatment at 500 mg·L⁻¹ rate showed lower values for %¹⁵Ndff than those at 250 mg·L⁻¹ rate for all leaf categories. The %¹⁵Ndff for 500 mg·L⁻¹ treatment values are (5.63, 5.66, 5.00) for the upper, medium and lower leaves, respectively. The %¹⁵Ndff tends to decrease with increasing leaf age. The high %¹⁵Ndff in the young leaves could be due to leaves being at their maximum activity and transpiration stage ^[36].

In our previous work ^[40] we found that using the stem injection technique in olive (cv. Frantoio) tree, the maximum increase in %¹⁵Ndff was observed at 200 mg·L⁻¹ boron rate for both leaves and flowers. In the current study, we found that boron has resulted in more translocation of nitrogen to the upper leaves of mango trees particularly at 250 mg·L⁻¹ rate. Our results agree with Miley et al. ^[29] and El-Motaium et al. ^[40] who found that boron enhances the utilization of nitrogen. Following boron application leaf became the dominant sink for nitrogen in the off-year.

The ¹⁵N is translocated to the bud and fruit as shown by the %¹⁵Ndff values. Boron application resulted in ¹⁵N translocation to the buds as %¹⁵Ndff values for buds are 6.37, 11.09, and 8.71 for 0.00, 250, 500 mg·L⁻¹ boron application rate, respectively (**Table 4**). The maximum %¹⁵Ndff value was obtained at 250 mg·L⁻¹ boron treatment. The ¹⁵N was also translocated to the fruit, peel and pulp (**Table 5**) with %¹⁵Ndff values of 0.734 and 0.628 for peel and pulp, respectively. The labeled N in the leaf of the off-year represents 190% of the total labeled N accumulated in fruit + bud.

3.3 Influence of boron on ¹⁵N internal cycling in mango tree organs in the "off-year"

Boron affects the distribution of nitrogen between the tree organs (leaf, bud and fruit). When boron was applied at 250 mg·L⁻¹ rate higher %¹⁵Ndff and %N was observed in the leaf than in the bud or the fruit. The descending order of %¹⁵Ndff in mango tree organs at 250 mg·L⁻¹ boron application rate is as follows: average of leaf categories (12.58) > bud

(11.09) > fruit peel (0.734) > fruit pulp (0.628). Boron applied at 500 mg·L⁻¹ rate shows lower %¹⁵Ndff values than those at 250 mg·L⁻¹ rate.

The total nitrogen percent of the upper leaves increased as the boron application rate increased as follows: 1.10, 1.36, 1.56 for 0.00, 250, 500 $\rm mg\cdot L^{-1}$ boron treatments, respectively. A similar trend was observed in the other leaf categories. Fruit peel contains higher total nitrogen percent than the pulp and was greatest at 500 $\rm mg\cdot L^{-1}$ boron.

The application of boron at 250 mg·L⁻¹ in the off-year has modified ¹⁵N internal accumulation among mango tree organs. More nitrogen (¹⁵N) accumulated in the leaf than in the bud or fruit, as shown by the high %¹⁵Ndff values. There is a tendency for an increase in total nitrogen percent in all the tree organs (leaf, bud and fruit) as the boron application rate increases. This may be due to boron enhancement of nitrogen uptake and tissue concentration ^[30].

Nason and McElory [57] found that boron is in-

volved in nitrogen metabolism, in particular the synthesis of the nitrogen base "uracil" in nucleic acid, RNA ^[42]. In addition, boron is involved in nucleic acid and protein metabolism ^[11]. Nucleic acids are required for the stimulation of growth ^[58], protein synthesis, photosynthesis, fruit set and yield.

Nitrogen flows to places where there is high metabolic demand [36] and boron is involved in nitrogen and hormone metabolism [58]. In the off-year the leaf represents an organ of high metabolic demand; therefore, more ¹⁵N-labelled nitrogen flows to the leaf under boron application. Thus, we suggest that the high accumulation of nitrogen in the leaf followed by boron application has stimulated plant growth and formation of the flowering hormone, upon its translocation to the growing points it stimulates flower induction, fruit set and yield. Our suggestion is in the same line with the early literature of Reece et al. [59] who emphasized leaf effects and hormonal factors in flower formation.

Upper leaf			Medium le	Medium leaf			Lower leaf		
% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	
0.767	7.41	1.05	0.631	6.10	1.19	0.541	5.23	1.24	
Bud % ¹⁵ N a.e. Bud %No			Bud %Ndf	6Ndff		Bud %N			
0.65 6.28				1.23					
Fruit peel			-		Fruit pulp				
% ¹⁵ N a.e.		% ¹⁵ Ndff %N % ¹⁵ N a.e. % ¹⁵ Ndff		%N					
1.25	12.08 1.07 0.93		8.99		1.17				

Table 3. Leaf % ¹⁵N a.e., %Ndff and %N in the off-year.

Boron (mg·L ⁻¹)	Upper leaf			Medium leaf			Lower leaf		
	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N
0.0 (control)	0.542	5.24	1.10	0.449	4.34	1.02	0.400	3.86	1.00
250	1.442	13.93	1.36	1.340	12.95	1.13	1.124	10.86	1.28
500	0.583	5.63	1.56	0.586	5.66	1.52	0.517	5.00	1.44
LSD (0.05)	0.255	2.292	0.064	0.022	0.794	0.080	0.032	0.299	0.082

Table 4. Bud %15N a.e., %Ndff and %N in the off-year.

Boron (mg·L ⁻¹)	% 15N a.e.	% ¹⁵ Ndff	%N
(control)	0.659	6.37	1.0
250	1.148	11.09	1.18
500	0.902	8.71	1.33
LSD (0.05)	0.012	0.408	0.102

Table 5. Fruit %¹⁵N a.e., %Ndff and %N in the off-year.

Boron (mg·L ⁻¹)	Fruit peel			Fruit pulp	Fruit pulp		
	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	% ¹⁵ N a.e.	% ¹⁵ Ndff	%N	
0.0 (control)	0.045	0.435	1.19	0.029	0.280	1.08	
250	0.076	0.734	1.25	0.065	0.628	1.17	
500	0.056	0.541	1.31	0.045	0.435	1.24	
LSD (0.05)	0.0044	0.0023	0.035	0.0021	0.0013	0.044	

3.4 Comparison between "on and off-year"

The on-year trees exhibited a greater $\%^{15}$ Ndff than the off-year trees likely as a result of N dilution with growth. The buds accumulated almost similar percent of ($\%^{15}$ Ndff = 6.28) in the on-year and in the off-year ($\%^{15}$ Ndff = 6.37), at zero boron rate (**Tables 2 and 4**). With boron application, at 250 mg·L⁻¹ rate in the off year, the buds accumulated higher 15 N ($\%^{15}$ Ndff = 11.09) than the buds of the on-year ($\%^{15}$ Ndff = 6.28).

Results in **Tables 2 and 3** show that on year, upper, middle and lower leaves contain higher (%¹⁵Ndff) than those of the off year, at zero boron rate (control treatment). The on-year upper leaf contains %¹⁵Ndff = 7.41 while the off-year contains %¹⁵Ndff = 5.24. The descending order of %¹⁵Ndff was the same in the on and off years (upper leaves) > (middle leaves) > (lower leaves). However, the off-year showed lower values than the on-year. This indicates that the accumulation and utilization of N maintained the same trend in both years [on-year (2 months post N-15 injection)].

Comparing the accumulation of ¹⁵N in the on and off years, there is a greater accumulation of nitrogen in fruits of the on year than of those of the off-year. This could be due to the greater reproductive demand for nitrogen in the on year than in the off year. Our results agree with Weinbaum et al. ^[32]. Boron

application in the off-year at 250 mg·L⁻¹ resulted in twice as much ¹⁵N accumulation in the leaf than that of the on-year.

3.5 Effect of boron foliar spray in the off-year on yield, flowering and fruit set

In the on-year, fruit yield averaged 67.4 kg/tree while in the off-year yield in the control (zero B) trees was 9 kg/tree. Boron foliar application at 250 mg·L⁻¹ rate in the off-year caused a significant increase in fruit yield (33.5 kg/tree) relative to the control (9.0 kg/tree). The percent increase in mango fruit yield due to boron was 372% relative to the control and alternate bearing was alleviated by 37% ^[24]. This indicates a synergistic relationship between B and N.

Flower and final fruit set percentage significantly increased in response to boron foliar application (**Table 6**). The highest flower percentage and final fruit set percentage relative to the control was observed at 250 mg·L⁻¹ boron rate. Our results agree with Hegazi et al. [41] who found that boron rate of (200 mg·L⁻¹) coincides with the highest flower percentage, fruit set and yield in olive cv. Frantoio. Along the same line, Negi et al. [2] found an increase in mango (cv. Dashehari) fruit set as a result of boron foliar application at 200 ppm. Boron function in increasing nucleic acid and hormone synthesis [11] could stimulate growth, flowering, fruit set and yield in the off-year.

B concentration (mg·L ⁻¹)	Yield (kg/tree)	Flowering (%)	Final fruit set (%)
0.00	9.0	18.6	27.0
250	33.5	33.0	87.0
500	24.0	21.5	41.0
LSD (0.05)	0.79	0.79	8.8

Table 6. Effect of boron application rates on mango tree yield, flowering and final fruit set in the off-year.

4. Conclusions

We concluded that the highest uptake and translocation of ¹⁵N in the on and off years was achieved by the upper (young leaves) compared with the middle and lower leaves. In the on-year, fruits maintained the highest %Ndff. While in the "off-year", buds maintained the highest %¹⁵Ndff without boron application. However, boron application at 250 mg·L⁻¹ rate in the off-year resulted in higher ¹⁵N enrichment in the leaf, bud and fruit.

In the "on-year" the distribution pattern of the labeled nitrogen (¹⁵N) tended to show more enrichment in the fruit than in the other organs (leaf, bud). This result indicates that fruit has a greater reproductive demand for nitrogen than the other tree's organs. In the "off-year", the distribution pattern of the labeled nitrogen (¹⁵N) tends to exhibit more enrichment in the bud, without boron application (control treatment). However, after boron application, at 250 mg·L⁻¹ rate, the distribution pattern of ¹⁵N was altered in favor of the leaf.

Foliar boron application in the off-year enhanced nitrogen translocation and accumulation in all tree organs (leaf, bud, fruit). This indicates a functional association between boron and nitrogen. There is evidence of a synergistic relationship (positive interaction) between boron and nitrogen (boron facilitates N uptake, translocation and accumulation). This was achieved at a boron rate of (250 mg·L⁻¹) which coincides with the highest flowering, fruit set and yield. The results prove that our hypothesis is true.

Author Contributions

First author contribution: Develop the research idea and objective, execution of the experiments,

conduct the statistical analysis and writing of the manuscript.

Second author contribution: Provide help in the development of the research concept and review the manuscript.

Third author contribution: Provide help in the development of the research concept and review the manuscript.

Fourth author contribution: Provide help in conducting the experimental part of the research.

Conflict of Interest

The authors declare that there is no conflict of interest.

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