

Melia volkensii の遺伝的能力とプラスツリー特性表 (概要)

ケニア森林研究所 (KEFRI) と JICA は、東アフリカの乾燥・半乾燥地域 (ASAL) の社会林業に関し 35 年以上にわたる協力を進め、これまでに実施したプロジェクトにおいて地域への受け入れられやすさを基本として樹種を選定してきた。その精査した樹種の中から、*Melia volkensii* (以下、メリア) は今後の展開に最も適した樹種とりわけ乾燥地での商業樹種として優先づけられた。メリアは、家畜飼料、薬、養蜂の蜜源に用いられるなど用途が広く、さらに成長が速く高品質な木材が得られることで高く評価される郷土樹種である。直近 2 回のプロジェクトでは、このメリアの育種改良を進めてきた。

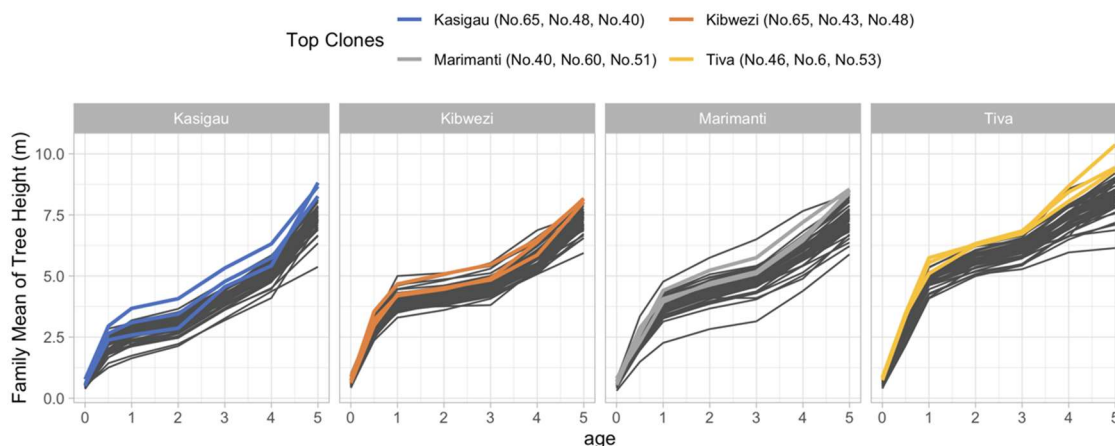
ケニアの乾燥地・半乾燥地からメリアのプラス木 (CPTs) の選抜を 2012 年から 2013 年にかけて行った。国内各地の天然分布域より、成長、幹の通直性、枝の大きさ、健全性等を基準とし、計 100 個体を選定した。

プラス木のクローンを植栽した採種園を 2013 年から Kitui および Kibwezi に設定した。設定に当たっては、プラス木から採取した穂を KEFRI Kitui 苗畑で 2012 年 9 月につぎ木によりクローン苗として育成した上で、採取園にプラス木あたりクローン苗 5 本、100 プラス木全体で 500 本のブロックを 6 反復、計 3000 本植栽した。

プラス木の遺伝的能力の評価及び次世代優良個体の選抜のため、次代検定林の設定を行った。採種園で得られた自然交配種子 (採種園で一本も種子が得られなかったプラス木クローンについては原木から採種した自然交配種子) を用い、プラス木の子供にあたる実生個体を育成し植栽した。メリアの天然分布域および気候区分を基準として区分された 4 つの生態地域ごとにメイン検定林とサブ検定林を設定し計 8 検定林とした。

検定林において成長等の表現型の計測を行った。樹高、直径を植栽 1 ヶ月後および 6 ヶ月後から 5 年後まで年 2 回測定した (コロナによる中断期間は除く)。幹の通直性は 5 段階の指数で、病害耐性は 2 段階で、着花および結実による繁殖性は 4 段階で評価した。成長の良いクローンは、成長の良い検定林では平均樹高 10m を越えるものも確認された。

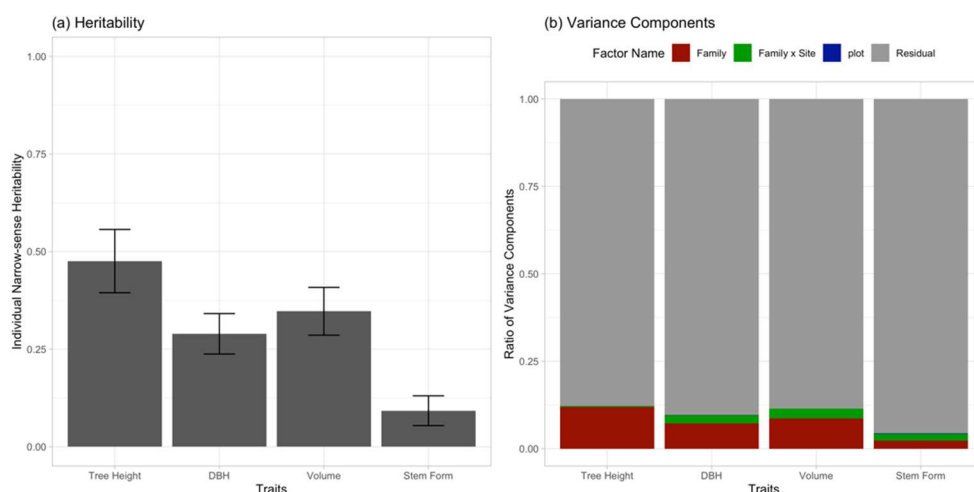
(a) Tree Height



図：メイン 4 検定林における樹高のクローン平均値の推移 (n=55)

メイン検定林の5年後計測データを用いて、遺伝性の評価および育種価の推定を行った。成長の遺伝的分析のため被害木（先折れ、病虫害など）や異常値を除き、55本のプラス木の自然交配実生2,664個体を評価対象とした。

遺伝性の評価の結果、メリアの成長性および幹の通直性は選抜による遺伝的改良を行うのに十分な遺伝性があることが確認された。4箇所の試験地を通した樹高の遺伝率は0.48であり、他の造林樹種と変わらない遺伝性であった。また、プラス木の遺伝的能力の発揮の程度が、植栽環境によりプラス木ごとに異なる程度を示す指標（下図(b)の緑部分の比率）は特に成長についてはそれほど大きいものではなく、採種園産種苗が当面は広い環境範囲で利用可能である可能性が示された。



図：形質ごとの遺伝率 (a)および分散成分 (b)

各プラス木の遺伝的能力を示す値である育種価を推定し、それに基づいて採種園の改良のシミュレーションを行った。その結果、採種園の上位50%の個体を用いて種子を生産することにより、植栽後5年時の樹高で約7%、材積で約17%の向上が見込まれることが予測された。

表：採種園の上位50%を選抜した場合の改良効果

Trait	Genetic Gain	General Mean	Relative Genetic Gain
Tree Height	0.54 m	7.56 m	7.13 %
DBH	0.75 cm	11.9 cm	6.29 %
Volume	0.0082 m ³	0.0490 m ³	16.77 %
Stem Form	0.12	3.52	3.33 %

各プラス木の育種価を、特性表としてまとめた。この特性表は、劣勢なプラス木クローンの間伐を通じた採種園産種子の改良や、メリアの次世代育種における選抜に貢献し、最終的にはケニアにおけるメリアのより生産的な商業植林の造成につながると期待される。

表：全検定林を通して評価したプラス木特性表（上位 30 クローンを抜粋）

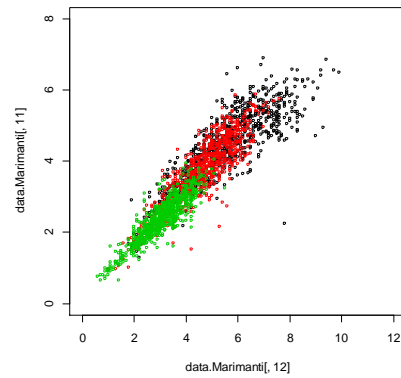
Clone	Stem Volume*		Tree Height*		DBH*		Stem Form*		Fruits**	Fungal Disease**	Assessed Trees***
	m ³	rank	m	rank	cm	rank	index	rank	rate	rate	number
No. 6	0.0246	1	1.300	1	1.680	3	0.351	1	0.10	0.07	57
No. 9	0.0210	2	0.748	9	1.924	2	0.318	2	0.13	0.01	111
No. 7	0.0199	3	0.655	13	2.069	1	0.210	7	0.10	0.07	52
No. 43	0.0173	4	0.686	11	1.475	4	0.115	13	0.12	0.16	21
No. 49	0.0172	5	0.768	8	1.423	5	0.250	3	0.13	0.04	45
No. 51	0.0153	6	1.209	3	0.994	9	0.026	24	0.18	0.16	36
No. 48	0.0130	7	1.232	2	0.778	14	-0.045	33	0.00	0.17	18
No. 29	0.0124	8	0.379	17	1.307	7	0.237	6	0.19	0.02	46
No. 53	0.0109	9	0.939	5	0.813	11	-0.049	35	0.10	0.29	20
No. 11	0.0108	10	0.322	21	1.366	6	0.096	16	0.22	0.01	100
No. 3	0.0100	11	0.650	14	0.825	10	0.247	5	0.07	0.01	107
No. 40	0.0094	12	0.772	7	0.655	15	0.132	11	0.18	0.13	48
No. 57	0.0082	13	0.380	16	1.057	8	-0.004	28	0.03	0.03	29
No. 46	0.0080	14	0.674	12	0.559	18	-0.050	36	0.00	0.21	11
No. 4	0.0063	15	0.249	22	0.790	13	0.184	8	0.17	0.04	118
No. 8	0.0061	16	-0.139	31	0.813	12	-0.174	49	0.08	0.01	96
No. 60	0.0054	17	0.883	6	0.180	23	-0.203	52	0.11	0.22	14
No. 19	0.0043	18	0.112	26	0.560	17	-0.047	34	0.16	0.11	54
No. 20	0.0039	19	-0.234	35	0.617	16	0.124	12	0.18	0.11	54
No. 22	0.0029	20	0.690	10	0.003	28	0.027	23	0.15	0.04	53
No. 14	0.0028	21	0.146	25	0.205	22	0.183	9	0.20	0.10	44
No. 58	0.0026	22	0.159	24	0.358	21	-0.042	32	0.08	0.17	10
No. 33	0.0022	23	-0.221	34	0.490	19	-0.120	41	0.07	0.18	46
No. 16	0.0018	24	0.206	23	0.138	25	0.052	18	0.26	0.04	55
No. 65	0.0011	25	0.340	20	0.100	26	0.042	20	0.09	0.00	11
No. 5	0.0008	26	0.365	19	-0.322	34	0.180	10	0.10	0.07	83
No. 2	0.0008	27	0.068	27	-0.060	30	0.112	14	0.14	0.01	101
No. 70	0.0005	28	0.437	15	-0.330	35	-0.075	38	0.04	0.13	42
No. 74	0.0004	29	-0.094	30	0.165	24	-0.119	40	0.09	0.09	48
No. 61	0.0002	30	0.953	4	-0.613	39	0.111	15	0.16	0.05	41

*: The breeding value, presented as the deviation from average. **: The rate of fruits bearing or fungal infected tree to the total investigated trees across 4 main-sites. ***: investigated number of trees for growth traits across 4 main-sites.

Genetic Performance and Plus Tree Traits Table for *Melia volkensii* in the Drylands of Kenya

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Foreword

Kenya Forestry Research Institute (KEFRI) has been collaborating with the Japan International Cooperation Agency (JICA) in a series of projects more than 35 years related to social forestry development in the ASAL areas of eastern Kenya. From the projects implemented so far, appropriate tree species for the region have been identified based on their adaptability and acceptance by the local communities. Among the species screened, *Melia volkensii* was prioritized as the most suitable for further development especially for commercial forestry in the drylands. *M. volkensii* is an indigenous tree species that is highly rated for its fast growth and multiple uses. The species is used for production of high quality timber, fodder, medicine and as bee forage.

In 2012, a project titled “Project on development of drought tolerant trees for adaptation to climate change in drylands of Kenya (2012~2017)” was initiated and has been implemented with the aim of enhancing KEFRI’s research capacity in breeding of drought tolerant trees. In 2016 to 2021, another project “Capacity Development Project for Sustainable Forest Management in Kenya” was implemented. This project continued with some of the tree breeding activities initiated by the project on “Development of drought tolerant trees”

During implementation of the two projects, KEFRI collected candidate plus trees of *M. volkensii* and evaluated their genetic performance for growth, stem form, flower and fruits bearing and health status through establishment of seed orchards, establishment of progeny tests and their evaluation for the last six years. The genetic performance of the plus trees has been evaluated and the results presented in this report as Plus Tree Traits Table. The traits table summarizes growth, reproductive and disease tolerance parameters across the test sites. The Plus tree traits table will contribute to orchard improvement through rogueing of inferior families leading to more improved seed from the orchards. The results will also guide selection for further breeding programs in subsequent *Melia* breeding generations ultimately leading to establishment of more productive *Melia* commercial plantations in Kenya.

Joshua K Cheboiwo

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1. Introduction

In Kenya, the arid and semi-arid areas (ASALs) occupy 80% of the country's land area and are home to over 10 million people who are predominantly pastoralists and agro-pastoralists. The drylands are endowed with abundant natural resources, with more than 70% of national livestock population, and 90% of the wild game that supports much of the tourism sector. The population pressure on drylands is caused by immigration of farming communities from the high potential areas, changing lifestyles of the local inhabitants, bioprospecting and minerals exploration. Consequently, drylands are increasingly being recognized as the new frontier for Kenya's development.

The climatic conditions in the drylands are generally hot, with low, erratic and poorly distributed rainfall. Annual rainfall ranges from 150 mm to 750 mm and occurs in one or two peaks around April and November. The potential evapotranspiration is higher than precipitation. Effects of climate change, which is a permanent shift in the traditional climate patterns, are also expected to exacerbate the already harsh climatic conditions in drylands. It is projected that in the next 100 years, the average temperature in the East Africa region could increase by 3 °C, as a result of climate change. The frequency and intensity of extreme climatic events such as droughts and floods will also increase, and the arid and semi-arid lands (ASALs) will be most affected.

Kenya is more susceptible to impacts of climate change due to frequent and recurrent droughts, over dependence on rain-fed agriculture and the limited economic capacity. The effects of climate change are already being felt in Kenya and the Government has given official recognition of the current and potential impacts of climate change through the preparation of a policy on disaster management and also in the Vision 2030. Among the goals of the Vision include; formulating adaptation programs on climate change and desertification in ASALs. The National Forest Program 2016-2030 recommends expansion of commercial trees in ASALs and establishment of tree improvement programme for the drylands.

In the past 34 years, the Japan International Cooperation Agency (JICA) has collaborated with Kenya Forestry Research Institute (KEFRI) and Kenya Forest Service (KFS) in social forestry development in the ASAL areas of eastern Kenya. From the projects implemented so far, the appropriate tree species for the region have been identified based on their adaptability and acceptance by the local communities. Among these, *Melia volkensii* was prioritized as the most suitable for further development. *M. volkensii* is an indigenous tree species that is highly rated for its fast growth and multiple uses. The species is used for production of high quality timber, fodder, medicine and as bee forage.

In 2012, a project titled "Project on development of drought tolerant trees for adaptation to climate change in drylands of Kenya (2012~2017)" was initiated and has been implemented

with the aim of enhancing KEFRIs research capacity in breeding of drought tolerant trees for Kenya's drylands and development of an extension system to promote planting of indigenous trees in ASALs. Furthermore, KEFRI had implemented "Social Forestry for Adaptation to Climate Change (2014 to 2018)" as a third country training program, which has been supported by JICA since 1995. From 2016 to 2021, another project "Capacity Development Project for Sustainable Forest Management in Kenya" was implemented. This project was a continuation of the breeding project for development of drought tolerant trees.

The objective of tree breeding is to supply genetically superior materials that are adapted to target planting areas, solve specific problems and produce desired products. To achieve this objective, tree breeding is mainly consists of "determining variation patterns, assessing the intensity and cause of variation, packaging the variation into desirable trees, and mass production of the improved individuals" (Zobel and Talbert, 1984).

In the first project on breeding for drought tolerance (2012-2017), KEFRI selected 100 Candidate Plus Trees (CPTs) of *M. volkensii*, collected their propagules and established seed orchards. Progeny trials to evaluate genetic worth of the CPTs in terms of growth, flower and fruits bearing and health status were also established but evaluated in the second project. In this report, we compile results of analysis of the genetic performance of plus trees and present them as traits table.

The results of genetic analyses and the resulting traits table will contribute to orchard improvement through roguing of inferior families leading to more improved seed from the orchards. The results will also guide selection for further breeding programs in subsequent *Melia* breeding generations ultimately leading to establishment of more productive *Melia* commercial plantations in Kenya. We provide brief description for the materials, measurement of traits, and analysis methods used before presenting the plus-tree traits table.

2. Selection of *Melia* Candidate Plus Trees and establishment of clonal seed orchards

2.1 Candidate Plus Tree selection

One hundred (100) Candidate Plus Trees (CPTs) of *Melia volkensii* were selected from 2012 to 2013 across arid and semi-arid areas the Kenya. The CPTs were selected in various regions within the species' natural range using a set of criteria (Table 1). The locations where CPTs were selected is depicted in Figure 1. The selection process involved reconnaissance in sites of natural occurrence of the species, followed by actual selection within the viable *Melia* populations (at least 30 individuals). Individual CPTs selection involved identification of potential candidate tree, assessment of the selected tree and its five nearest neighbours (check trees). The purpose of assessing the candidate *M. volkensii* tree and the check trees was to confirm superiority of the plus tree in comparison with the neighbouring *Melia* trees. In addition to documentation of geographic location of the CPTs (GPS), vegetation type, soil type and climatic conditions of the site was done. Candidate plus tree selection was based

on stem straightness, growth vigor, branch size and number and health (Table 1).

Candidate Plus Trees were selected in the following regions (transects): Mutha-Inyali, Katulani-kavisuni, Voi-Mwatate, Voi-Galana, Embu-Ishiara-Gatunga, Embu-Dams, Mwea Special, Mwingi-Nuu, Mwingi-Tseikuru, Isiolo-Meru, Garissa-Bangale, Garba-Wamba and Wamba-Marsabit.

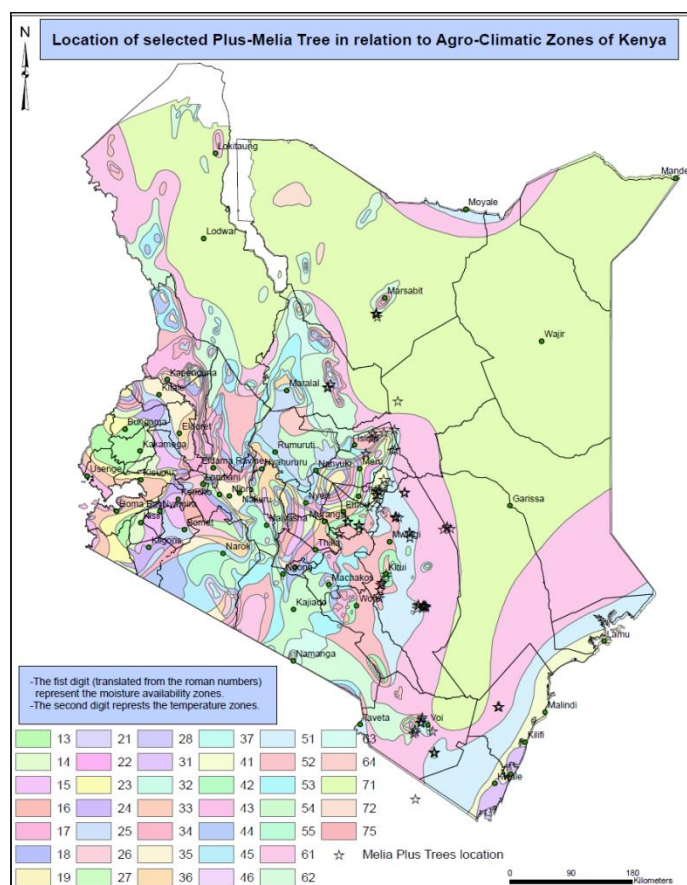


Figure 1: *Melia volkensii* Plus tree selection sites in Kenya

Table 1: Selection Criteria for *Melia volkensii* CPTs

No	Criteria
1	A tree in the dominant or co-dominant crown class (at or above the general tree canopy level) compared with the surrounding trees;
2	Superior in height and diameter growth in comparison to surrounding 5 trees
3	High growth vigour in comparison to surrounding 5 trees
4	Good tree form (Straight straightness and light branching habit)*
5	Light-medium branching, less steep angled branches
6	Not crooked or twisted stems/branches, No spiral grain
7	Free from insect pests and free of any signs of diseases

2.2 Establishment of seed orchards

Rootstocks of *M. volkensii* were raised at the KEFRI Kitui nursery for 4 months. Scions from the 100 CPTs trees were collected and taken to the nursery for grafting onto the rootstocks. Eighty (80) scions were obtained from each of the selected plus trees and 72 grafted onto Melia rootstocks in September 2012 using the top grafting method. The grafted seedlings were raised and managed in the nursery through watering, root pruning and disease control for 4 months until early December. The seed orchards were established from 2013 at Kitui and Kibwezi, using the grafted seedlings. In each orchard, 100 clones of plus trees each with 5 ramets were planted in 6 blocks resulting to 3,000 seedlings per site. The spacing was 6 m x 6 m (Figure 2).

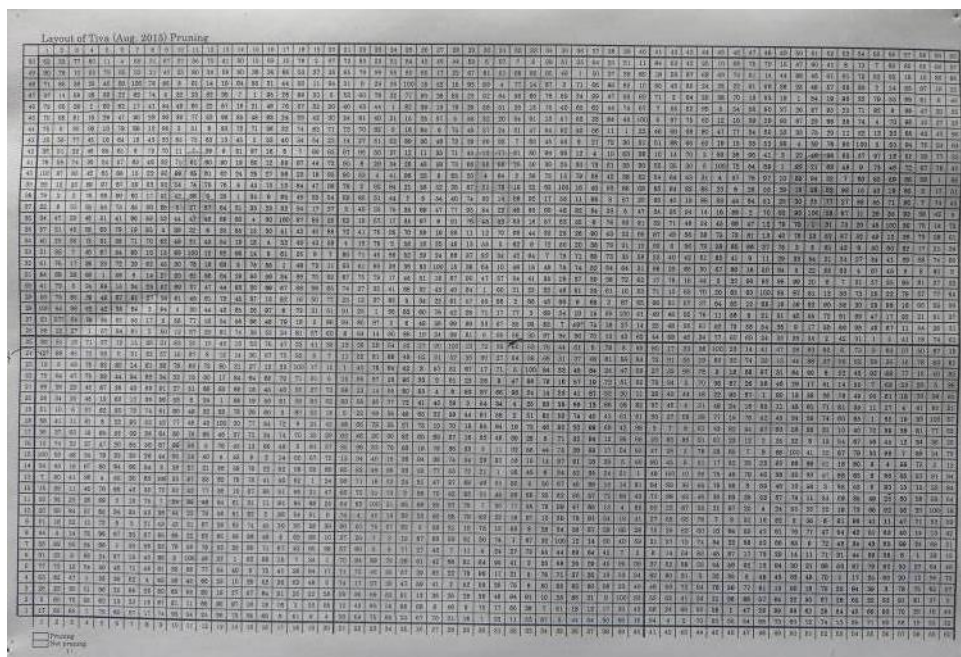


Figure 2: Layout of Melia Orchard in Tiva, Kitui

3. Establishment of *Melia volkensii* progeny tests

Seed for establishment of progeny tests was collected from the seed orchards, and from the original CPTs in the field for orchard families that had not started seeding. The seedlings were raised at KEFRI Kitui nursery in 2015 and used to establish 4 main progeny and 4 sub-progeny tests.

The eight progeny tests were located in four eco-regions (wet and cool (in the north), semi-wet and semi-cool (middle), semi-dry and semi-hot (middle), and dry and hot (south)) based on climate and the distribution range of *M. volkensii* (Figure 3). The progeny tests were of two types, main and sub-progeny tests, located in Marimanti (main) and Gaciongong (sub), Makima (sub) in the North; Tiva (main) and Kibwezi (main) and Ikithuki (sub) in the middle areas and Kasigau (main) and Voi (sub) in the south.

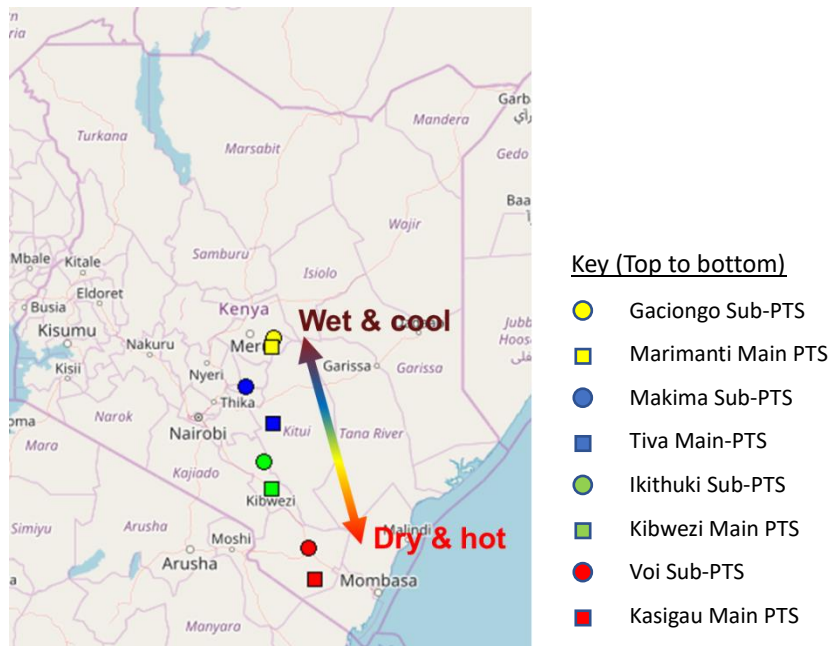


Figure 3. Climatic range in locations of progeny tests
(The dots represent progeny test sites across the region)

4. Phenotypic and genetic analysis of growth, tree form, fecundity and health at progeny trials

Tree height, Diameter at Ground Level (DGL) and Diameter at 50cm (D_{50}) were assessed one month after planting. Tree height, Diameter at Breast Height (DBH) and D_{50} were assessed twice a year starting at 6 months to age 5 years. Height was measured using a measuring rod while diameter was assessed using calipers and diameter tape. Stem straightness was evaluated using a subjective score rating of a 5-point scale: 1 = most twisted, 5 = straightest starting at age 4 years. Tolerance to fungal disease were measured with a subjective rating with a 2-point scale: 1 = no disease, 2 = diseased starting at 4 years. Fecundity was also assessed at year on a 4-point scale 1-None, 2-flowers, 3-fruits and 4-fruits and flowers starting at age 4 years. The analysis was done using data from the main progeny tests.

4.1 General growth trends

The average growth for height of trees at about 5 years ranged from 3.4 m to 11.2 m for Marimanti, 3.9 m to 11.4 m for Tiva, 3.8 m to 9.7 m for Kibwezi and 3.7 m to 9.8 m for Kasigau. In diameter growth the values ranged from 3.1 cm to 20.8 cm for Marimanti, 4.1 cm to 21.6 cm for Tiva, 3.1 cm to 18.1 cm for Kibwezi and 4.6 cm to 17.7 cm for Kasigau. However, due to wide variation, the mean values trends per site are shown in Figures 4 (height) and 5 (diameter).

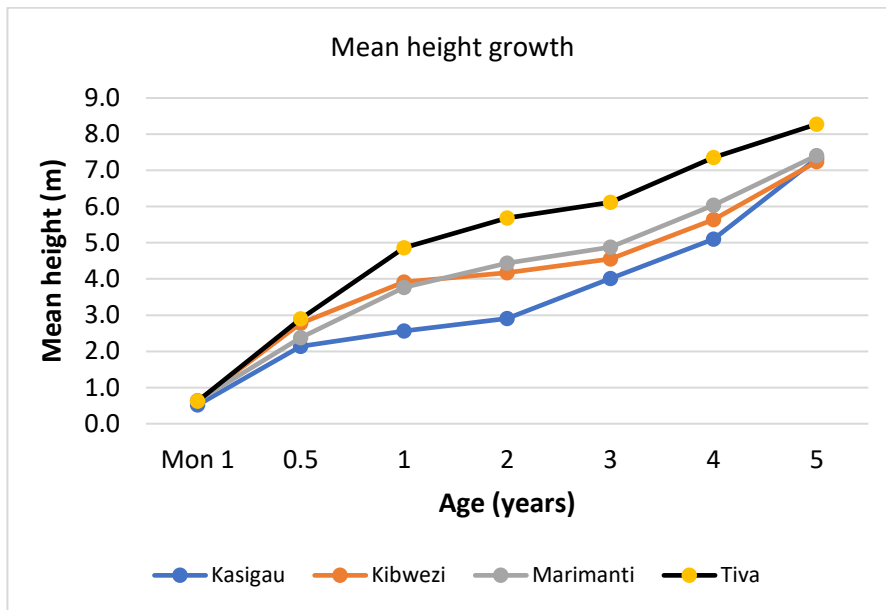


Figure 4. Mean height of *Melia volkensii* at 4 progeny test sites

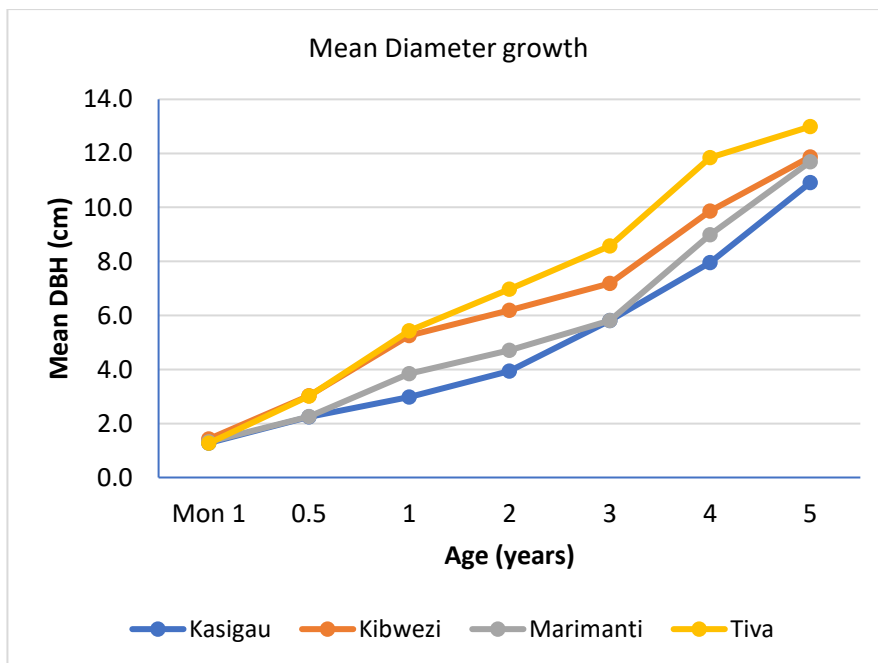


Figure 5. Mean diameter of *Melia volkensii* at 4 progeny test sites

4.2 Best performing clones by site in terms of tree height and DBH

The family height and DBH mean of each clone at each site is shown in Figures 6a and 6b. The best performing clones in height and DBH were different among sites and traits are shown on top of Figures 6a and 6b.

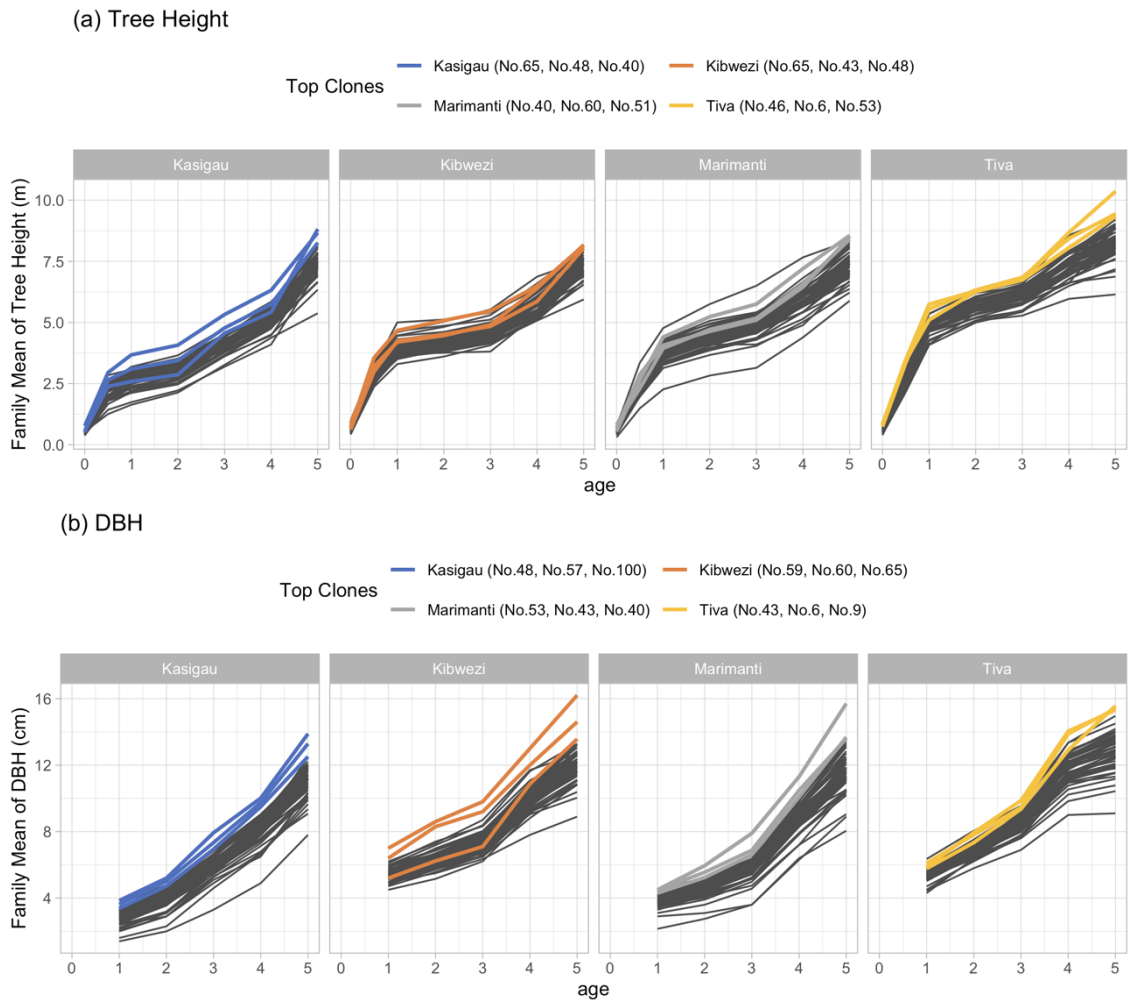


Figure 6. Family mean growth of *Melia volkensii* for each clone at 4 PT sites
(Top 3 clones at age 5 for each site were presented with color.)

4.3 Calculation of stem volume

Since there are no stem volume equations available for *M. volkensii*, we used stem volume equation for *M. azedarach* in Japan (Forestry Agency Japan, 1970) as shown below:

$$V = 10^{-4.199274} \times DBH^{1.8007256} \times H^{0.98511581}$$

Where V is stem volume of a tree (m³), DBH is diameter at breast height (cm), and H is tree height (m).

The volume obtained from above equations fall in the middle of conic volume and cylindrical volume calculated from tree height and DBH (Figure 7). Since the two species are in the same genus and exhibit similar characteristics, usage of this equation is appropriate until an original stem volume equation is constructed for *M. volkensii* in future. Stem volumes were calculated using this equation in subsequent analysis.

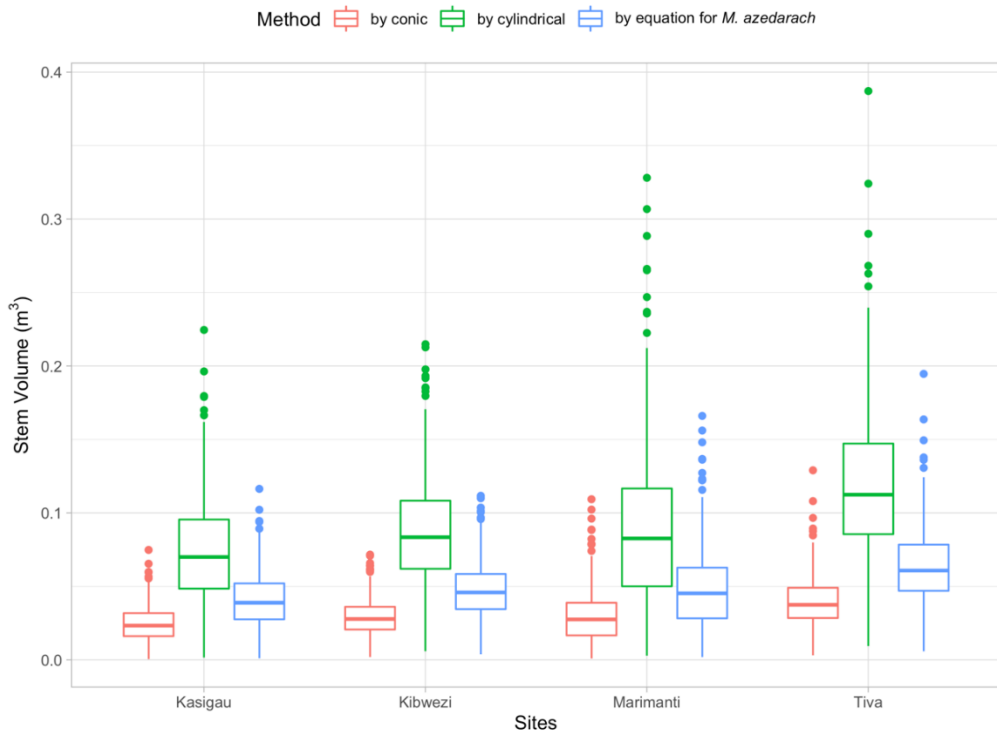


Figure 7. The difference in stem volume among progeny test sites and different calculation methods

(The boxplot shows individual stem volumes at age 5).

The difference of family mean of stem volume increased with age. The top clones were different among sites. The best performing clones in terms of stem volume in Kasigau were clone No 46, 77 and 65, in Kibwezi clones No. 59, 60 and 65, in Marimanti clone no 40, 43 and 53 and in Tiva clone no 6, 43 and 46 (Figure 8).

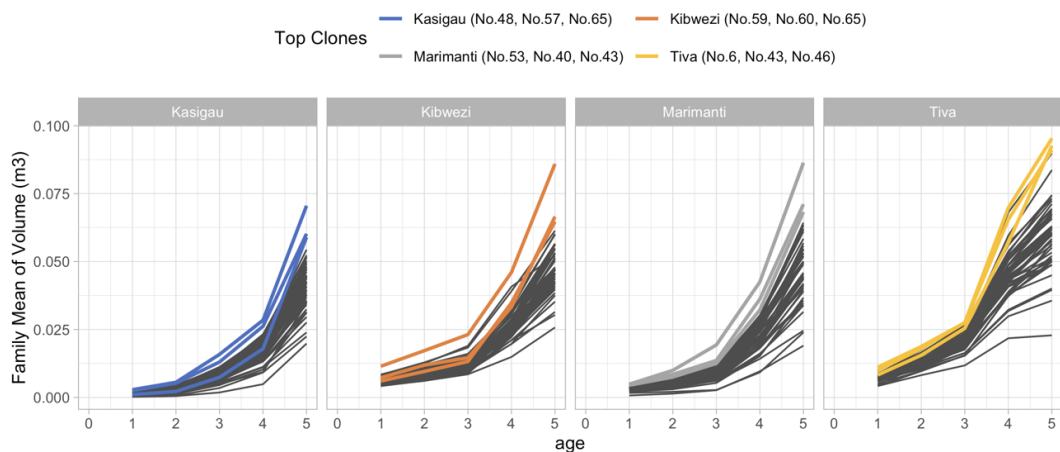


Figure 8. Family mean stem volume of *Melia volkensii* for each clone at 4 progeny test sites

(Top 3 clones at age 5 for each site were presented with color)

The performance shown above contains site effect. However, selection of clones for breeding 2nd and subsequent generations will be subject to the ranking presented in plus tree traits table (Section 6). The top 3 families presented per site should not be taken as recommendation for planting in the areas.

5. Genetic analysis of phenotyped traits

5.1 Data cleaning

We used the 4 main progeny test sites data for analysis, because the number of planted progeny trees for each plus trees were small (5 to 1, mostly 2) in sub-progeny sites. For genetic analysis of growth data at age 5, data collected from damaged trees (broken top, infected, and so on) were removed. The outliers were checked by tree height-DBH ratio and the data which the ratio was > 2 or the ratio < 0.2 were removed. The numbers of trees are different among mother plus trees.

Progenies of 69 out of the collected 100 CPTs were planted in the main progeny test sites. After data cleaning, the available growth data were from 2,664 progeny trees of 55 plus trees planted in four main progeny tests were used in the analysis.

5.2 Evaluation of fecundity and disease tolerance

The number of progeny trees which flowered were 46 which translated to 1.3% of the total trees in 4 main PTS at age 5 years. The number of trees with fruits were 477 representing 13.5% of total trees in 4 main PTS at age 5 years. The fruits bearing rate of the progenies of each CPT for each site, and their relationship among sites are presented in Figure 9. Even between Kibwezi and Marimanti, where relatively higher fruits bearing rates were observed, the correlation coefficient was low ($r = 0.193$).

The number of progeny trees which were infected by fungal disease were 240 representing 6.8% of all trees in 4 main PTS at age 5 years. The rate of infected trees for the progenies of each plus-tree at each main site, and their relationship among sites are presented in Figure 10. The infection rates were biased to zero, and the relationship is weak among sites.

The genetic parameters and breeding values were not estimated for fruit bearing and infection rates because of their zero biased distribution and low relationship among PTS sites. The rate for each clone calculated for all trees across 4 main-sites were used for the evaluation value for each plus tree.

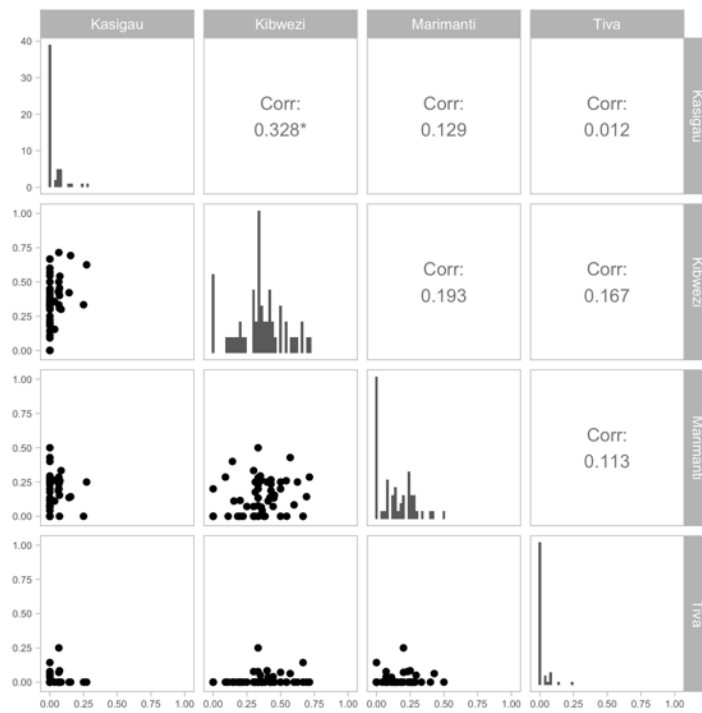


Figure 9. The relationship of fruits bearing rate of the open-pollinated progenies of plus trees among sites.

(Points in lower triangle panel shows the fruits bearing progeny tree rate of each plus tree).

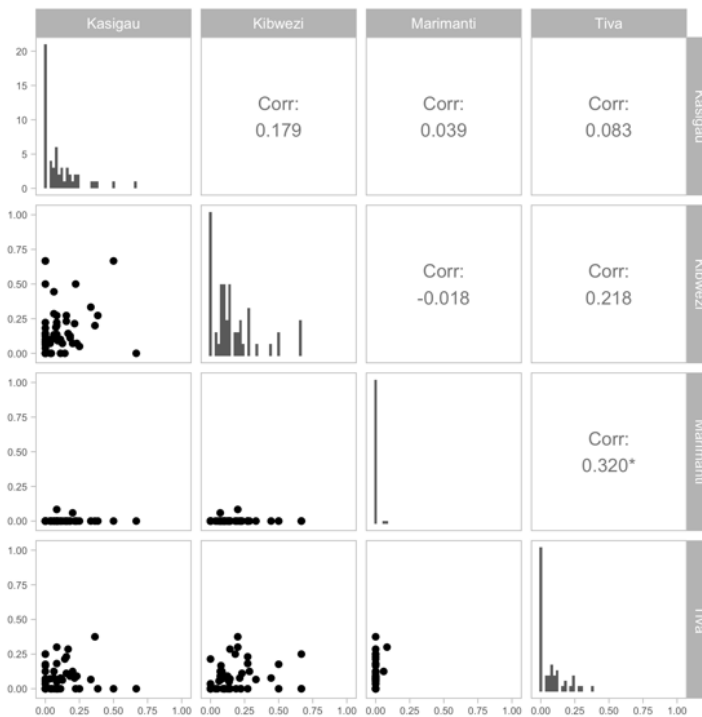


Figure 10. Rate of plus tree infections among sites.

(Points in lower triangle panel show the rate of infection among progenies of each plus tree).

5.3 Genetic parameters for growth and tree form traits at each site and genetic correlation among sites.

Genetic parameters were analyzed for tree height, DBH, stem volume, and tree form at age 5 for each PTS site, using following mixed linear model:

$$y_{ijk} = \mu + B_i + F_j + p_{ij} + e_{ijk}$$

where y_{ijk} is the phenotypic value for target trait, μ is the mean, B_i is the fixed effect of replication i , F_j is the random effect of mother plus tree (general combining ability (GCA) of mother clone) j , p_{ij} is random effect of plot by replication i and clone j , and e_{ijk} is random residual effects. Tree form traits were treated as interval scale in this analysis whereas they were measured on ordinal scale. Variance components were estimated from the linear model by REML method, and heritability for each site and trait were obtained using following equation:

$$h^2 = \frac{4 \cdot \sigma_f^2}{\sigma_f^2 + \sigma_p^2 + \sigma_e^2}$$

Where h^2 is individual narrow-sense heritability, σ_f^2 is family variance, σ_p^2 is plot variance, and σ_e^2 is residual variance. The BLUP of GCA of mother tree clones were also obtained using above mixed linear model. The breeding value of each CPT was calculated as twice the GCA for each mother plus tree for each site. The genetic correlations among sites were calculated as correlation coefficients of breeding values of CPTs per site.

The heritability values are shown in Figure 11. Tree height showed highest heritability. The heritability of stem form was low, but showed that tree form is a heritable trait in *M. volkensii*. Tiva, where average growth was best, showed the highest heritability compared to other sites.

The genetic correlations among sites for tree height and stem volume are presented in Figure 12. All combinations showed positive relationships. The genetic correlation of tree height between Tiva and Marimanti showed highest genetic correlation ($r = 0.602$). Tree height showed higher genetic correlations than stem volume in most combination of sites.

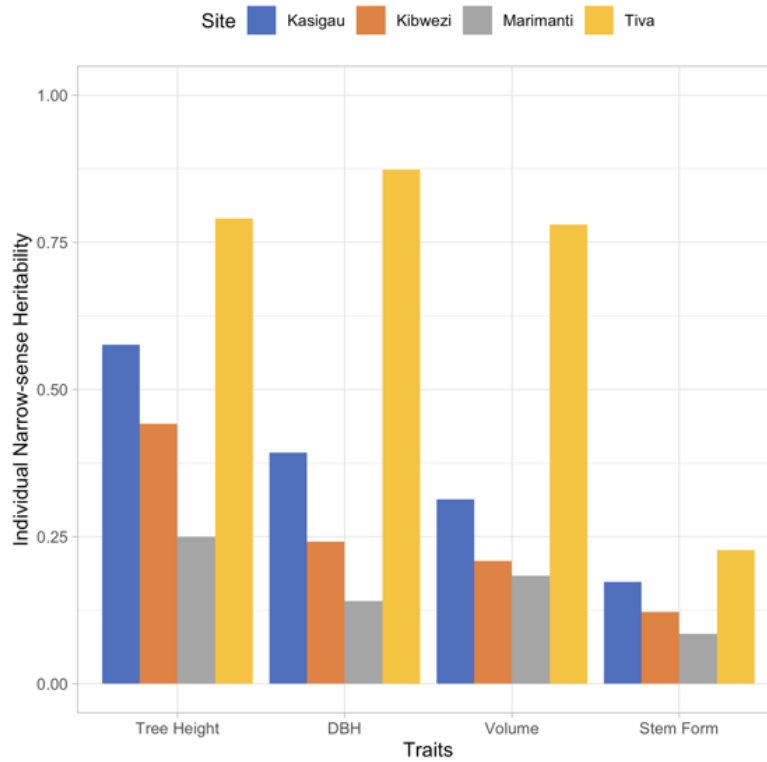


Figure 11. The narrow sense heritability of growth and tree at age 5

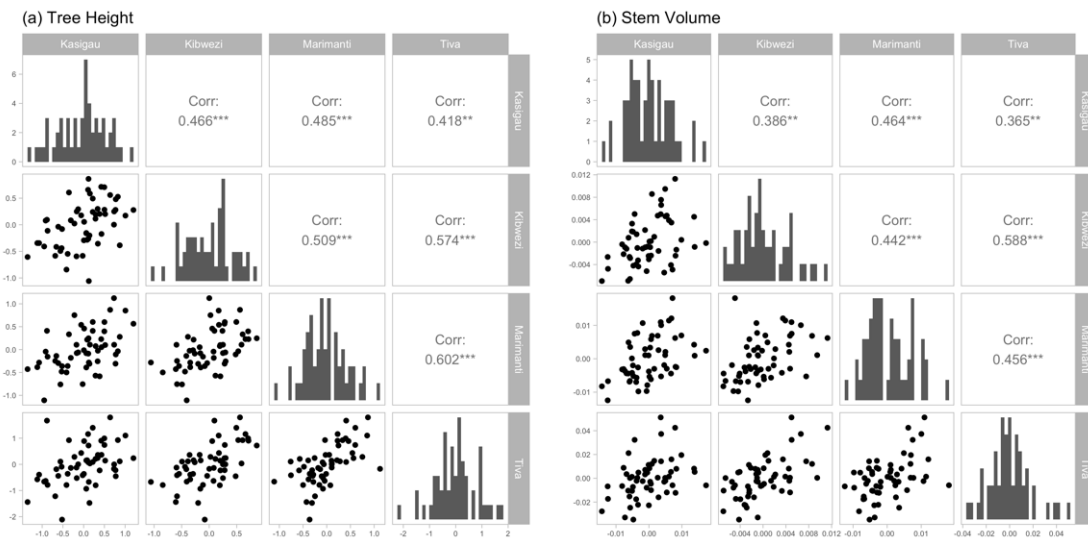


Figure 12. The genetic correlations of tree height and stem volume among sites.

(Each point shows the breeding value of CPT at each site at age 5 yrs.).

5.4 The effect of seed source on performance the traits

There are two seed sources, from seed orchard and from the original trees in the field, as described in section 3. The effects of seed source for each site and each trait were tested at age 5 years. We prepared data set for each site that have clones including both seed sources.

The number of clones included in tests were 22.6 on average. A log likelihood ratio test was adopted for the two linear models as explained in 5.4. The effect of seed source was added to the model as fixed effects. The p-values were over 0.05 for all traits and sites. This indicates that the effects of seed source for the tested traits in *M. volkensii* were small and non-significant and therefore were not included in subsequent analysis.

5.5 Prediction of breeding values across sites

Genetic parameters were analyzed for tree height, DBH, stem volume, and tree form at age 5 years across sites, using following mixed linear model:

$$y_{ijkl} = \mu + E_i + EB_{ij} + F_k + EF_{ik} + p_{ijk} + e_{ijkl}$$

Where y_{ijkl} is the phenotypic value for target trait, μ is mean, E_i is fixed effect of site i , EB_{ij} is the fixed effect of replication j in site i , F_k is the random effect of mother plus tree k (general combining ability (GCA) of CPT), EF_{ik} is the effect of interaction between site i and mother clone k (genotype by environment interaction), p_{ijk} is random effect of plot by replication j and clone k at site i , and e_{ijkl} is random residual effect. The individual narrow sense heritabilities were estimated using the following equation:

$$h^2 = \frac{4 \cdot \sigma_f^2}{\sigma_f^2 + \sigma_{sf}^2 + \sigma_p^2 + \sigma_e^2}$$

Where h^2 is individual narrow-sense heritability, σ_f^2 is family variance, σ_{sf}^2 is family by site variance, σ_p^2 is plot variance, and σ_e^2 the residual variance. The breeding value of each CPT was calculated as twice the GCA for each CPT per site. The analysis was executed using breedR package (Muñoz and Sanchez 2020) in statistical package R (R Core Team 2021).

The narrow sense heritabilities are shown in Figure 13(a). Tree height showed highest heritabilities (0.476) compared to other assessed traits. The heritability of stem form was low, but genetic improvement would be possible. The ratios of variance components are shown in Figure 13 (b). The variance components of family by site interaction in tree height was small, showing small effects of environment on ranking of tree height across the sites. The effects of family by site interaction were larger for DBH and stem volume compared to tree height. There is a possibility that some families are more suited to different eco-region based on genetic correlation among sites (Figure 12) and the ratio of family by site interaction variance to family variance (Figure 13). However, the selection of superior plus tree groups adapted for all eco-region would be possible. Several progeny sites would be required for each eco-region to clarify the effects of genotype by eco-region interaction in the *M. volkensii* breeding program.

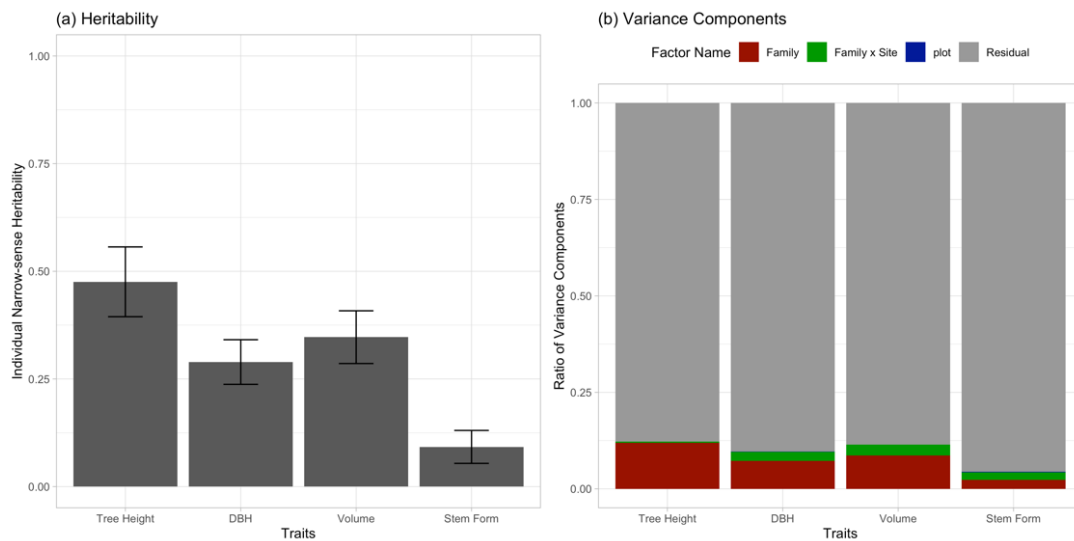


Figure 13. Heritabilities and variance components estimated across main progeny test sites

The relationship among breeding value, growth traits, tree form, ratio of fruits bearing trees, and ratio of disease infected trees are presented in Figure 14. Stem form showed higher positive correlation of around 0.6 with growth traits – height and DBH. The correlation among the ratio of fruits bearing trees and growth traits, and among disease infected rate and growth rate was low.

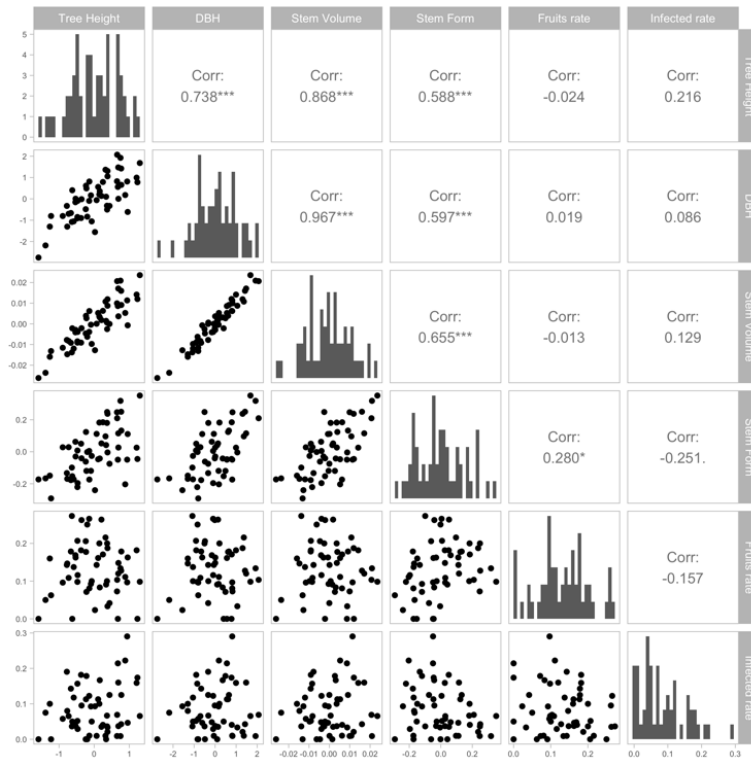


Figure 14: The relationship among breeding value of growth traits and tree form, across-sites fruits bearing rate and infection rate.

The relationship between breeding values predicted at each site and breeding values predicted across sites were shown in Figure 15 for tree height and stem volume. The correlation coefficients were higher for tree height. The correlations were high, but in some site showed smaller values under 0.7. This means the selected clones based on breeding values predicted across sites have the possibility that shows smaller performance than expected.

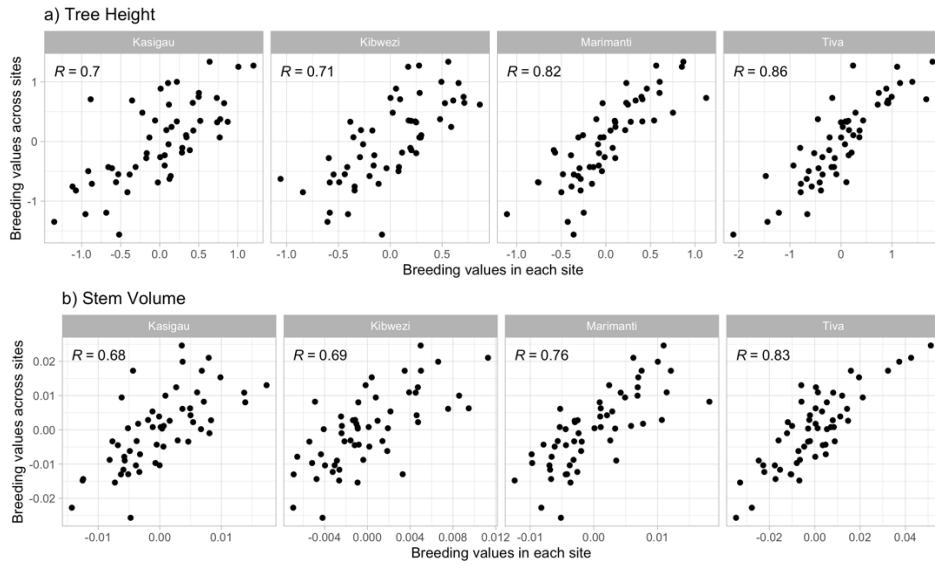


Figure 15: The relationship between breeding values predicted in each site and breeding values predicted across sites.

(a): Tree height, b): Stem volume.

5.6 Simulation of genetic improvement of seed orchards

The expected genetic gain by selection of superior clones and rouging inferior ones from seed orchards were calculated based on breeding values for each trait:

$$G_a = \frac{\sum_{i=1}^m B_k}{m} - \frac{\sum_{i=1}^n B_k}{n}$$

Where G_a is genetic gain, m is number of selected plus trees, n is number of pre-selected plus trees, B_k is breeding values of k the clone. B_k is ordered to descending order of breeding values. The selection rate is m/n . The relative gain was calculated by:

$$G_r = \frac{G_a}{M}$$

Where G_r is relative gain, M is general mean (average of fixed effects across sites). The assumptions were: random mating among all trees, no pollen contamination from outside sources, and equal contribution to pollen and seed production among all clones in a seed orchard. The genetic gain by selection of top 50% of clones for each trait are shown in Table 2. By rouging seed orchards by 50% of inferior clones, the stem volume in the stand at age 5 years will be improved by about 0.0082 m³, and its rate to general mean is about 16%. This

shows that stem volume in plantations would be improved by about 16 % if the seed is obtained from rogued seed orchards compared to if the seed is from orchards with all 55 plus trees. For tree height, the genetic gain is 0.54 m, and will be improved by about 7%.

The genetic gains by changing selection rate are presented in Figure 16. When top 16% of clones (top 9 clones) are selected, volume would be improved about 33% at stand age 5. Because the genetic correlations are positive among traits (Figure 12), when the superior clones are selected based on the breeding value of volume, tree form would be also improved to some extent.

Table 2. Genetic gain by the selection of top 50 % of clones age 5 across PTS sites

Trait	Genetic Gain	General Mean	Relative Genetic Gain
Tree Height	0.54 m	7.56 m	7.13 %
DBH	0.75 cm	11.9 cm	6.29 %
Volume	0.0082 m ³	0.0490 m ³	16.77 %
Stem Form	0.12	3.52	3.33 %

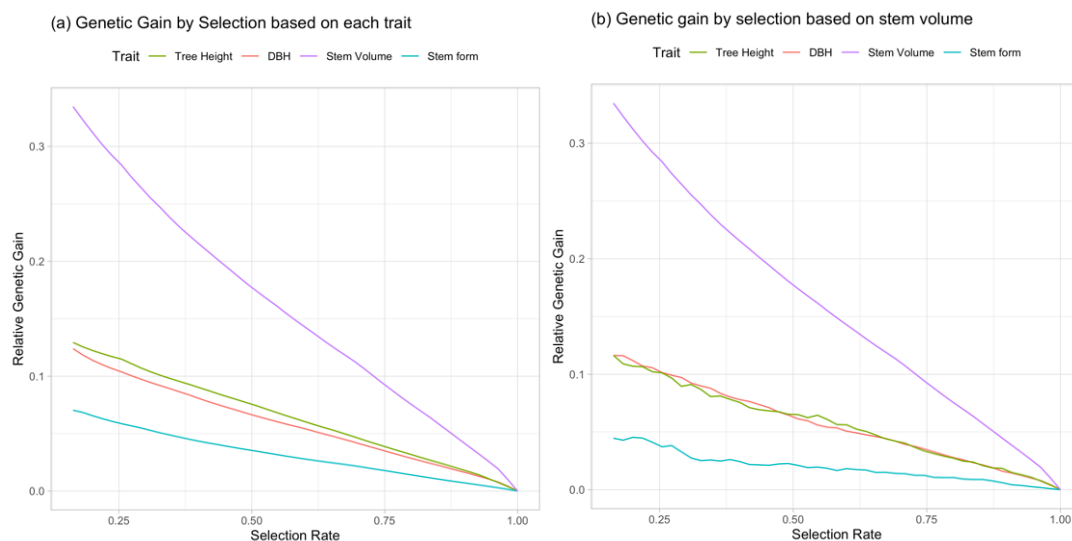


Figure 16. Genetic gain with different selection rates.

(Selection rate means the rate of selected clones. (a) Selections were based on each trait. (b) Selections were based on stem volume.)

6. Plus tree traits table

The plus tree traits across all sites are presented in Table 3. The number of CPTs listed in the Table are 55 clones, and ordered by the deviations from average breeding values of stem volume. Data obtained from 4 main CPTs at age 5 was used. The calculation of breeding values is described in section 5.5. The plus tree trait table that shows breeding values ordered by the deviations from the average for each site are presented in Table 4.

This plus trees traits table will contribute to orchard improvement through rogueing of inferior families leading to more improved seed from the orchards. After deciding on the rogueing intensity (%) to be used the orchards, the table can be used to determine which families are to be removed. Similarly, it can be used in determining families to select forward for second and subsequent breeding generations.

Table 3. Plus Tree traits table across sites

Clone	Stem Volume*		Tree Height*		DBH*		Stem Form*		Fruits**	Fungal Disease**	Assessed Trees***
	m ³	rank	m	rank	cm	rank	index	rank	rate	rate	number
No. 6	0.0246	1	1.300	1	1.680	3	0.351	1	0.10	0.07	57
No. 9	0.0210	2	0.748	9	1.924	2	0.318	2	0.13	0.01	111
No. 7	0.0199	3	0.655	13	2.069	1	0.210	7	0.10	0.07	52
No. 43	0.0173	4	0.686	11	1.475	4	0.115	13	0.12	0.16	21
No. 49	0.0172	5	0.768	8	1.423	5	0.250	3	0.13	0.04	45
No. 51	0.0153	6	1.209	3	0.994	9	0.026	24	0.18	0.16	36
No. 48	0.0130	7	1.232	2	0.778	14	-0.045	33	0.00	0.17	18
No. 29	0.0124	8	0.379	17	1.307	7	0.237	6	0.19	0.02	46
No. 53	0.0109	9	0.939	5	0.813	11	-0.049	35	0.10	0.29	20
No. 11	0.0108	10	0.322	21	1.366	6	0.096	16	0.22	0.01	100
No. 3	0.0100	11	0.650	14	0.825	10	0.247	5	0.07	0.01	107
No. 40	0.0094	12	0.772	7	0.655	15	0.132	11	0.18	0.13	48
No. 57	0.0082	13	0.380	16	1.057	8	-0.004	28	0.03	0.03	29
No. 46	0.0080	14	0.674	12	0.559	18	-0.050	36	0.00	0.21	11
No. 4	0.0063	15	0.249	22	0.790	13	0.184	8	0.17	0.04	118
No. 8	0.0061	16	-0.139	31	0.813	12	-0.174	49	0.08	0.01	96
No. 60	0.0054	17	0.883	6	0.180	23	-0.203	52	0.11	0.22	14
No. 19	0.0043	18	0.112	26	0.560	17	-0.047	34	0.16	0.11	54
No. 20	0.0039	19	-0.234	35	0.617	16	0.124	12	0.18	0.11	54
No. 22	0.0029	20	0.690	10	0.003	28	0.027	23	0.15	0.04	53
No. 14	0.0028	21	0.146	25	0.205	22	0.183	9	0.20	0.10	44
No. 58	0.0026	22	0.159	24	0.358	21	-0.042	32	0.08	0.17	10
No. 33	0.0022	23	-0.221	34	0.490	19	-0.120	41	0.07	0.18	46
No. 16	0.0018	24	0.206	23	0.138	25	0.052	18	0.26	0.04	55
No. 65	0.0011	25	0.340	20	0.100	26	0.042	20	0.09	0.00	11
No. 5	0.0008	26	0.365	19	-0.322	34	0.180	10	0.10	0.07	83
No. 2	0.0008	27	0.068	27	-0.060	30	0.112	14	0.14	0.01	101
No. 70	0.0005	28	0.437	15	-0.330	35	-0.075	38	0.04	0.13	42
No. 74	0.0004	29	-0.094	30	0.165	24	-0.119	40	0.09	0.09	48
No. 61	0.0002	30	0.953	4	-0.613	39	0.111	15	0.16	0.05	41
No. 31	-0.0010	31	-0.361	37	-0.015	29	-0.041	31	0.26	0.10	37
No. 100	-0.0031	32	-0.624	45	0.404	20	-0.153	44	0.00	0.05	38
No. 13	-0.0033	33	0.369	18	-0.503	38	0.249	4	0.20	0.04	53
No. 10	-0.0034	34	-0.529	42	0.008	27	-0.030	30	0.16	0.04	76
No. 17	-0.0034	35	-0.144	32	-0.094	32	0.005	26	0.27	0.12	41
No. 34	-0.0043	36	-0.471	39	-0.093	31	-0.196	51	0.12	0.03	66
No. 27	-0.0045	37	-0.535	43	-0.157	33	0.062	17	0.17	0.05	77
No. 1	-0.0049	38	-0.241	36	-0.484	37	-0.062	37	0.21	0.00	33
No. 37	-0.0062	39	-0.198	33	-0.836	44	-0.002	27	0.25	0.00	8
No. 23	-0.0071	40	-0.779	50	-0.331	36	-0.133	42	0.14	0.19	33
No. 55	-0.0079	41	-0.728	48	-0.701	41	0.029	21	0.11	0.07	42

Clone	Stem Volume*		Tree Height*		DBH*		Stem Form*		Fruits**	Fungal Disease**	Assessed Trees***
	m ³	rank	m	rank	cm	rank	index	rank	rate	rate	number
No. 32	-0.0088	42	-0.477	41	-0.851	45	-0.221	53	0.10	0.16	42
No. 76	-0.0090	43	-0.477	40	-0.910	47	0.008	25	0.21	0.05	55
No. 59	-0.0097	44	-0.374	38	-0.900	46	0.050	19	0.18	0.18	9
No. 12	-0.0104	45	-0.664	47	-0.664	40	-0.177	50	0.18	0.14	71
No. 21	-0.0104	46	-0.072	29	-1.056	49	-0.026	29	0.10	0.05	56
No. 30	-0.0117	47	-0.897	51	-0.804	43	0.028	22	0.10	0.06	48
No. 41	-0.0123	48	-0.583	44	-1.055	48	-0.172	47	0.14	0.12	45
No. 91	-0.0130	49	0.022	28	-1.556	53	-0.240	54	0.02	0.09	37
No. 97	-0.0130	50	-1.229	52	-0.803	42	-0.292	55	0.06	0.00	16
No. 39	-0.0144	51	-0.636	46	-1.081	50	-0.099	39	0.27	0.04	52
No. 36	-0.0148	52	-0.768	49	-1.301	52	-0.169	46	0.15	0.08	69
No. 24	-0.0154	53	-1.264	53	-1.301	51	-0.149	43	0.16	0.10	44
No. 67	-0.0227	54	-1.387	54	-2.184	54	-0.166	45	0.05	0.08	36
No. 79	-0.0257	55	-1.587	55	-2.751	55	-0.173	48	0.00	0.00	9

*: The breeding value, presented as the deviation from average. **: The rate of fruits bearing or fungal infected tree to the total investigated trees across 4 main-sites. ***: investigated number of trees for growth traits across 4 main-sites.

Table 4: Plus Tree traits table for each site

clone	Across Sites**		Kasigau***						Kibwezi***					
	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank
No. 6	0.0246	1	0.639	9	0.250	22	0.0036	18	0.559	7	0.537	10	0.0050	7
No. 9	0.0210	2	0.498	12	1.262	5	0.0079	7	0.707	3	1.237	2	0.0113	1
No. 7	0.0199	3	0.118	25	0.731	12	0.0037	16	0.863	1	0.758	5	0.0066	5
No. 43	0.0173	4	-0.354	40	-0.482	39	-0.0043	39	0.607	5	0.596	8	0.0050	6
No. 49	0.0172	5	0.501	11	0.866	11	0.0068	9	0.283	13	0.470	13	0.0035	13
No. 51	0.0153	6	1.007	2	1.071	6	0.0098	4	0.172	23	-0.037	30	0.0004	21
No. 48	0.0130	7	1.200	1	2.196	1	0.0173	1	0.277	15	-0.133	34	-0.0002	24
No. 53	0.0124	8	0.106	27	0.273	21	0.0027	20	0.658	4	0.481	12	0.0047	9
No. 29	0.0109	9	0.219	21	0.867	10	0.0061	11	0.241	18	0.620	7	0.0039	12
No. 11	0.0108	10	0.735	7	1.926	3	0.0137	3	0.249	17	0.705	6	0.0045	11
No. 3	0.0100	11	0.423	14	0.040	26	0.0009	22	0.713	2	0.919	4	0.0086	3
No. 46	0.0094	12	-0.885	50	-0.682	42	-0.0062	47	0.095	25	0.110	22	0.0008	20
No. 40	0.0082	13	0.731	8	0.645	13	0.0071	8	0.002	30	-0.720	48	-0.0049	50
No. 57	0.0080	14	0.519	10	2.151	2	0.0139	2	0.200	20	-0.155	35	-0.0009	27
No. 4	0.0063	15	0.150	22	0.897	9	0.0049	14	0.587	6	1.259	1	0.0095	2
No. 8	0.0061	16	0.385	15	0.570	15	0.0037	17	0.203	19	1.129	3	0.0075	4
No. 60	0.0054	17	0.011	31	-0.120	30	-0.0012	32	0.055	28	0.353	16	0.0022	15
No. 19	0.0043	18	0.768	6	0.454	18	0.0050	13	0.280	14	0.578	9	0.0046	10
No. 58	0.0039	19	0.085	28	0.018	28	-0.0001	28	-0.283	39	-0.233	38	-0.0025	39
No. 20	0.0029	20	-0.173	37	-0.286	34	-0.0019	33	-0.594	52	0.047	25	-0.0012	32
No. 22	0.0028	21	0.824	4	0.987	8	0.0083	5	0.527	8	-0.433	40	-0.0014	34
No. 14	0.0026	22	0.337	17	0.085	24	0.0016	21	0.298	11	0.089	24	0.0009	19
No. 65	0.0022	23	0.774	5	0.631	14	0.0054	12	0.480	10	0.513	11	0.0047	8
No. 33	0.0018	24	0.006	32	-0.370	35	-0.0035	36	-0.293	40	-0.038	31	-0.0010	31
No. 70	0.0011	25	-0.223	38	0.546	16	0.0006	23	0.023	29	-0.526	44	-0.0024	37
No. 74	0.0008	26	0.112	26	0.154	23	0.0001	26	-0.263	38	0.264	18	0.0001	23

clone	Across Sites**		Kasigau***						Kibwezi***					
	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank
No. 16	0.0008	27	0.430	13	-0.161	32	-0.0011	31	-0.173	36	0.011	27	-0.0009	30
No. 61	0.0005	28	0.219	20	-1.260	52	-0.0052	43	0.493	9	0.097	23	0.0019	16
No. 5	0.0004	29	-0.060	34	-0.123	31	0.0004	25	0.177	22	-0.439	41	-0.0009	28
No. 2	0.0002	30	0.342	16	1.030	7	0.0067	10	-0.356	43	-0.736	51	-0.0041	46
No. 100	-0.0010	31	0.140	23	1.602	4	0.0081	6	-0.201	37	-0.296	39	-0.0024	38
No. 31	-0.0031	32	0.063	30	0.482	17	0.0029	19	-0.161	35	-0.182	36	-0.0016	35
No. 34	-0.0033	33	-0.612	46	-0.377	36	-0.0038	37	-0.037	31	0.440	15	0.0014	18
No. 10	-0.0034	34	-0.531	44	-1.251	51	-0.0077	51	-0.439	47	-0.008	28	-0.0021	36
No. 13	-0.0034	35	0.870	3	0.330	19	0.0046	15	-0.386	44	-0.933	53	-0.0054	52
No. 1	-0.0043	36	0.063	29	-0.234	33	-0.0005	29	-0.155	34	-0.034	29	-0.0008	26
No. 27	-0.0045	37	-0.386	41	-1.105	48	-0.0068	49	-0.547	49	0.306	17	-0.0012	33
No. 17	-0.0049	38	0.285	19	0.273	20	0.0006	24	0.113	24	0.134	20	0.0002	22
No. 37	-0.0062	39	-0.167	36	-1.149	49	-0.0049	41	0.250	16	0.232	19	0.0019	17
No. 55	-0.0071	40	-0.560	45	-0.443	37	-0.0032	34	-0.493	48	-0.725	49	-0.0043	48
No. 76	-0.0079	41	-0.659	47	-0.481	38	-0.0058	45	-0.415	46	-1.048	55	-0.0066	53
No. 23	-0.0088	42	-1.074	53	-0.848	45	-0.0081	52	-0.345	42	0.133	21	-0.0004	25
No. 32	-0.0090	43	-0.916	51	-1.245	50	-0.0057	44	0.079	27	-0.494	42	-0.0029	42
No. 12	-0.0097	44	-0.024	33	0.036	27	-0.0007	30	-0.584	51	-0.539	45	-0.0052	51
No. 21	-0.0104	45	0.289	18	0.061	25	0.0000	27	0.193	21	-0.499	43	-0.0031	43
No. 30	-0.0104	46	-0.414	42	-0.563	40	-0.0039	38	-0.843	54	-0.098	33	-0.0040	45
No. 91	-0.0117	47	-0.132	35	-0.821	44	-0.0059	46	0.293	12	-0.690	47	-0.0026	40
No. 97	-0.0123	48	-0.685	48	-0.051	29	-0.0033	35	-0.584	50	-0.233	37	-0.0032	44
No. 59	-0.0130	49	-0.309	39	-0.869	46	-0.0051	42	0.083	26	0.449	14	0.0033	14
No. 41	-0.0130	50	0.125	24	-0.927	47	-0.0063	48	-1.058	55	-0.961	54	-0.0069	54
No. 36	-0.0144	51	-1.117	54	-1.413	53	-0.0124	53	-0.344	41	-0.579	46	-0.0048	49
No. 24	-0.0148	52	-0.952	52	-1.709	54	-0.0125	54	-0.408	45	-0.077	32	-0.0026	41
No. 39	-0.0154	53	-0.870	49	-0.667	41	-0.0073	50	-0.113	33	0.025	26	-0.0009	29
No. 67	-0.0227	54	-1.352	55	-1.976	55	-0.0143	55	-0.606	53	-0.909	52	-0.0069	55
No. 79	-0.0257	55	-0.520	43	-0.820	43	-0.0047	40	-0.081	32	-0.736	50	-0.0042	47

clone	Across Sites**		Marimanti***						Tiva***					
	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank
No. 6	0.0246	1	0.869	2	0.696	7	0.0110	4	1.794	1	3.961	1	0.0512	1
No. 9	0.0210	2	0.399	12	0.595	11	0.0062	12	0.993	6	3.825	2	0.0425	2
No. 7	0.0199	3	0.247	15	0.961	5	0.0100	6	0.720	12	3.763	3	0.0373	3
No. 43	0.0173	4	0.334	13	0.601	9	0.0075	8	0.923	8	2.992	4	0.0323	4
No. 49	0.0172	5	0.602	6	0.974	4	0.0121	2	0.740	11	1.331	8	0.0159	7
No. 51	0.0153	6	0.853	3	0.442	14	0.0070	9	1.101	5	1.308	9	0.0195	6
No. 48	0.0130	7	0.565	7	0.027	24	0.0024	20	0.236	18	-1.180	44	-0.0059	36
No. 53	0.0124	8	0.230	16	0.561	12	0.0069	10	1.157	4	-0.618	37	0.0005	25
No. 29	0.0109	9	0.409	11	1.048	3	0.0114	3	0.089	25	0.291	26	0.0014	23
No. 11	0.0108	10	0.106	20	0.597	10	0.0042	15	-0.001	29	1.274	10	0.0081	14
No. 3	0.0100	11	0.229	17	0.666	8	0.0069	11	0.903	9	0.888	14	0.0121	10
No. 46	0.0094	12	0.410	10	0.399	15	0.0049	14	1.671	2	1.153	12	0.0212	5
No. 40	0.0082	13	1.125	1	1.318	1	0.0181	1	-0.171	33	-0.386	31	-0.0059	35

clone	Across Sites**		Marimanti***						Tiva***					
	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank	H (m)*	rank	DBH (cm)*	rank	V (m ³)*	rank
No. 57	0.0080	14	0.100	22	0.136	19	0.0009	25	0.142	22	0.751	16	0.0053	18
No. 4	0.0063	15	0.108	19	0.116	20	0.0011	23	0.035	27	0.336	24	0.0004	27
No. 8	0.0061	16	-0.584	52	-0.539	48	-0.0053	43	-0.231	36	2.026	5	0.0143	9
No. 60	0.0054	17	0.460	9	0.138	18	0.0029	18	0.888	10	-0.412	33	0.0032	22
No. 19	0.0043	18	-0.308	42	0.220	17	0.0021	21	-0.224	35	0.174	27	-0.0021	29
No. 58	0.0039	19	0.217	18	0.083	21	0.0010	24	0.157	20	1.220	11	0.0113	11
No. 20	0.0029	20	0.105	21	1.152	2	0.0107	5	-0.123	31	-0.033	28	-0.0024	31
No. 22	0.0028	21	-0.036	27	-0.477	44	-0.0025	30	0.927	7	0.439	22	0.0080	15
No. 14	0.0026	22	-0.256	38	-0.360	38	-0.0030	33	0.241	17	1.582	6	0.0147	8
No. 65	0.0022	23	-0.106	33	-0.248	35	-0.0030	32	-0.454	42	-1.072	43	-0.0119	44
No. 33	0.0018	24	-0.013	25	0.751	6	0.0077	7	-0.365	38	0.479	21	0.0004	26
No. 70	0.0011	25	0.752	4	-0.059	28	0.0058	13	0.285	16	-1.220	45	-0.0100	41
No. 74	0.0008	26	-0.088	32	-0.414	41	-0.0054	44	0.077	26	1.000	13	0.0084	13
No. 16	0.0008	27	0.052	24	0.031	23	0.0011	22	0.364	14	0.574	18	0.0071	16
No. 61	0.0005	28	0.603	5	-0.137	30	0.0001	26	1.398	3	-0.640	38	0.0013	24
No. 5	0.0004	29	0.541	8	-0.033	26	0.0033	17	0.434	13	-0.642	39	-0.0029	32
No. 2	0.0002	30	-0.025	26	-0.507	46	-0.0044	41	0.360	15	0.397	23	0.0054	17
No. 100	-0.0010	31	-0.311	43	-0.035	27	-0.0024	29	-1.475	54	-0.088	29	-0.0122	45
No. 31	-0.0031	32	-0.079	31	0.296	16	0.0026	19	-0.932	51	-1.358	46	-0.0160	47
No. 34	-0.0033	33	-0.296	41	-0.432	42	-0.0057	45	-0.441	41	0.294	25	-0.0021	30
No. 10	-0.0034	34	-0.481	49	0.063	22	-0.0031	34	-0.092	30	1.445	7	0.0088	12
No. 13	-0.0034	35	0.279	14	-0.298	36	-0.0019	27	0.115	23	-0.388	32	-0.0029	33
No. 1	-0.0043	36	-0.391	47	-0.696	51	-0.0079	51	0.150	21	0.546	19	0.0048	20
No. 27	-0.0045	37	-0.359	44	-0.351	37	-0.0044	39	-0.362	37	0.887	15	0.0035	21
No. 17	-0.0049	38	-0.569	51	-0.231	34	-0.0062	46	0.207	19	-0.594	36	-0.0047	34
No. 37	-0.0062	39	-0.077	30	-0.180	32	-0.0020	28	-0.525	43	-0.728	40	-0.0072	39
No. 55	-0.0071	40	-0.761	54	-0.648	49	-0.0098	54	0.106	24	0.660	17	0.0049	19
No. 76	-0.0079	41	-0.139	34	-0.832	54	-0.0066	47	-0.186	34	0.516	20	0.0002	28
No. 23	-0.0088	42	-0.283	40	0.001	25	-0.0032	35	-0.393	39	-0.476	34	-0.0065	37
No. 32	-0.0090	43	-0.044	28	0.517	13	0.0035	16	-0.633	45	-2.480	52	-0.0247	52
No. 12	-0.0097	44	-0.755	53	-0.726	52	-0.0097	53	-0.417	40	-0.560	35	-0.0079	40
No. 21	-0.0104	45	0.076	23	-0.395	40	-0.0038	37	-0.800	50	-2.588	53	-0.0227	51
No. 30	-0.0104	46	-0.499	50	-0.437	43	-0.0070	50	-0.790	49	-1.570	47	-0.0176	49
No. 91	-0.0117	47	-0.061	29	-0.812	53	-0.0069	49	0.002	28	-1.659	49	-0.0153	46
No. 97	-0.0123	48	-0.248	37	-0.107	29	-0.0026	31	-1.218	52	-2.313	50	-0.0223	50
No. 59	-0.0130	49	-0.181	36	-0.366	39	-0.0042	38	-0.137	32	-1.011	42	-0.0106	43
No. 41	-0.0130	50	-0.282	39	-0.218	33	-0.0044	40	-0.672	47	-0.809	41	-0.0104	42
No. 36	-0.0144	51	-0.385	46	-0.535	47	-0.0067	48	-0.575	44	-1.599	48	-0.0173	48
No. 24	-0.0148	52	-1.109	55	-0.976	55	-0.0124	55	-0.661	46	-0.150	30	-0.0068	38
No. 39	-0.0154	53	-0.155	35	-0.160	31	-0.0037	36	-0.786	48	-3.412	54	-0.0329	54
No. 67	-0.0227	54	-0.427	48	-0.680	50	-0.0083	52	-1.440	53	-2.383	51	-0.0278	53
No. 79	-0.0257	55	-0.362	45	-0.501	45	-0.0052	42	-2.111	55	-3.742	55	-0.0347	55

*The breeding value, presented as the deviation from average. **Breeding values and rank across 4 sites. The values are same as in table 3. ***Breeding values and rank at each site.

7. References

- Munoz, F. and Sanchez, L. 2020. breedR: Statistical methods for forest genetic resources analysts (R package version 0.12-5). <https://github.com/famuvie/breedR>
- Forestry Agency. 1970. Stand tree stem volume table of Kumamoto regional forestry office. Forestry Agency, Tokyo, Japan (in Japanese).
- R Core Team. 2021. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL <https://www.R-project.org/>.
- Zobel, B. and Talbert, J. 1984. Applied Forest Tree Improvement. The Blackburn Press, Caldwell.