

Estimation of Genetic Variability, Heritability and Genetic Advance of Some Wollega Coffee (*Coffea arabica* L.) Landrace in Western Ethiopia Using Quantitative Traits

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Abstract: Arabica coffee is the predominant commodity in contributing for foreign exchange in Ethiopia and improvement for yield and other desirable traits is highly momentous. Estimating genetic diversity is a prerequisite activity in plant breeding program for crop improvement. This study was designed to determine the extent of genetic variability among Wollega coffee landrace and importance of gene revealed in traits. The 26 genotypes were tested during the 2016/2017 cropping season at Mugi and Haru sub- enters using RCBD. The combined analysis manifested significant difference among genotypes only in node number per primary branch (NNPB), fruit traits and Coffee leaf rust (CLR) although significant difference recorded for 18 and 22 of the 23 traits at Haru and at Mugi, respectively. The difference between environments was significant for all traits, except for CLR, yield (YLD), leaf, some fruit and bean traits. Performance at Haru was less than at Mugi for all traits showed significant difference. Genotype x environment (GEI) was significant for all traits excluding NNPB, leaf length (LL), fruit width and CLR indicating inconsistency performance of Coffee genotypes. At Haru, high phenotypic coefficient of variation (PCV>20%) recorded for YLD (25.5%), CLR (110.0%) and number of secondary branch (NSB) (22.0%), but High genotypic coefficient of variation (GCV>20%) recorded only for CLR (99.6%). At Mugi, High PCV and GCV (>20%) recorded for YLD (38.6%) and CLR (98.4%). Heritability ranged from 10.0% (YLD) to 88.0% (BW) while genetic advance (GAM) ranged from 1.5% (LL) to 32.4% (NSB) at Haru. At Mugi, Heritability ranged between 31% (CLR) and 84.0% (bean thickness) and between 3.3% (LL) and 44.0% (YLD) for GAM. The present results elucidate the existence of moderate genetic diversity among genotypes for some traits at individual location indicating the possibility of improvement for desired traits via selection. For further diversity analysis, molecular characterization methods need to be carried out.

Keywords: *Coffea arabica* L., Genetic Advance, Genotypic Variance, Heritability, Phenotypic Variance

1. Introduction

Coffee belongs to the family *Rubiaceae* and the genus *Coffea* [1, 2], in which there are at least 141 species [3]. However; the only two of these species commercially used are Arabica coffee (*Coffea arabica* L.) providing 60% and Coffee canephora (*Coffea canephora* Pierre ex A. Froehner) providing 40% of world's production [4]. *C. arabica* L. is the only allotetraploid ($2n=4x=44$) and a predominantly self-pollinating species and *C. canephora* P. is a cross pollinating

diploid ($2n=2x=22$) species [5, 6].

Even though Arabica coffee grown and produced in different part of the World's countries, genetically diverse species exist in Ethiopia than anywhere else in the world. This enhanced botanists and scientists to agree that Ethiopia is the center of origin, diversification and dissemination of the *Coffea arabica* L. plant [7]. According to Labouisse and Bayetta [8], Ethiopia is considered as the diversification for Arabica coffee (*Coffea arabica* L.) and high genetic variability exist for yield and yield components, diseases and pest

resistance traits. Similarly, as study carried out by Mesfin and Bayetta [9] on Arabica coffee collection from Hararge and Abdulfeta [10] from Tepi, and another study by Olika et al. [11] on Limmu Coffea arabica L. collection using quantitative traits indicated the existence of high genetic diversity.

Although coffee is growing in different Ethiopian geography, it is produced in immense within specific agro-ecological zones and political boundaries in Ethiopia. Thus, North Zone (Amhara and Benishangul Gumuz), South West Zone (Wollega, Illubabor, Jimma-Limu, Kafa, Tepi and Bench Maji), Rift Zone (Rift North and Rift South), South East Zone (Sidamo, Yergacheffe, Bale and Central Eastern Highland) and Harar Zone (Arsi, East Hararge and West Hararge) are the five main coffee growing Zone areas in Ethiopia [12]. From these Zones, the main coffee producing areas of Ethiopia are found within the South West and South East. The presence of high environmental diversity, distinct variation in coffee quality within and between regions [13] and location specificity of improved varieties [14] forced breeder to evaluate the genetic divergence for each location by collecting from that area. This is crucial to release coffee variety of high yielder and maintaining typical quality for that area.

Despite Wollega is one of the potential Arabica coffee growing areas of Western Ethiopia, only four Arabica coffee varieties had been released from this areas' coffee gene pool (Haru-I, Cala, Menesibu and Sinde) by pure line variety development. The released varieties give lower yield as compared to varieties released from coffee landrace of south western Ethiopia. The major contributing factors for such low yield in Wollega include limited exploitation of the existing germplasm of the areas and lack of well characterized and distinctly variable breeding material that is readily available for breeding work [1]. This implies that knowledge of genetic diversity among elite breeding

materials and understanding the significance of gene in traits is important for yield improvement of the crop. Hence, the present study was carried out with the intention to estimate the extent of genetic variability, broad sense heritability and expected genetic advance of some Wollega Arabica coffee landrace based on yield and yield related traits for the next breeding program.

2. Materials and Methods

Description of Studying Areas

The experiment was conducted at Haru and Mugi agricultural research sub centers which are sub centers of Jimma agricultural research center. Mugi found in kellelem Wollega zone at 34° 00' to East and 8° 40' to North. It is 610 km far from Jimma city to North West direction. It is located at altitude of 1570 m a.s.l and receive 1655 mm annual rain fall. Also, it has Nitosol soil type [15] and minimum 17°C and maximum 29°C temperature for this location. Haru is located 35° 47' 56" to East and 8° 59' 21" to North, in West Wollega zone at altitude of 1752 m a.s.l. and 360 km away from Jimma city. The area receives annual rain fall of 1727 mm which is unimodal, the peak being July. In addition, it has an average maximum and minimum temperature of 27°C and 16°C respectively [16] and sand clay loam soil.

Experimental Materials, Design and Agronomic practice

The experiment was conducted during 2017/2018 cropping season, on 22 promising Wollega coffee accessions which were taken from different batch of base collections with four standards check (Table 1). RCBD design in three replications was used. The study was superimposed on the already established coffee planted in July 2015 with six plants per plot using spacing of 2 m by 2 m and 4 m between replications. All field management applied as recommended [17].

Table 1. Description of Wollega coffee Accessions background.

No.	Accessions	Woreda	Peasants Association	Collection altitude (m. a.s.l)
1	W02/98	Haru	Wora Baro	1740
2	W34/98	Haru	Wora Baro	1790
3	W98/98	Haru	Chageli	1800
4	W141/98	Gimbi	H. Giorgis	1620
5	W163/98	Gimbi	Homa Arsama	1600-1670
6	W167/98	Gimbi	Homa Arsama	1600-1670
7	W175/98	Gimbi	Homa Arsama	1600-1670
8	W188/98	Gimbi	Homa Biribir	1550-1600
9	W191/98	Gimbi	Homa Biribir	1500-1570
10	W203/98	Gimbi	Siba Yesus	1560
11	W212/98	Gimbi	Sibo Charo	1560
12	W01/99	Haru	Guracha Holata	1660
13	W40/99	Haru	Dogi Adere	1720
14	W109/99	Ayira Gliso	-	1600
15	W03/00	Ayira Guliso	Waro Seyo	1500
16	W09/00	Ayira Guliso	Boke Keda	1600
17	W50/00	Ayira Guliso	Kurfessa birbir	1580
18	W52/00	Ayira Guliso	Kurfessa birbir	1520
19	W06/01	Ayira Guliso	Lalo Asella	1600
20	W08/01	Ayira Guliso	Tosiyo mole	1620
21	W15/01	Ayira Guliso	Buro Hasabar	1700
22	W38/01	Ayira Guliso	Nebo Daleti	1600

Checks / Released varieties				
1	Mana sibu (W78/98)	Haru	Haru	1550
2	Sinde (W92/98)	Haru	Haru	1590
3	Chala (W76/98)	Haru	Haru	1740
4	Haru-I (66/98)	Haru	Haru	1800

Methods and Data Recorded

Three randomly selected plants from each plot were used to record the plant growth parameters. However; for yield and disease data all plants per plot were used to record the necessary data. Data were recorded following the IPGRI descriptor [18]. Data taken during the study were Plant height (PH) (cm), Height up to first primary branch (HFPB) (cm), Total node number of main stem (TNN), Internodes length of the main stem (IL) (cm), Diameter of the main stem (DM) (mm), Stem habit (SH), Number of primary branches (NPB), Number of secondary branches (NSB), Average length of primary branches (ALPB) (cm), Number of nodes per primary branch (NNPB), Number of bearing primary branches (NBPB), Percentage of bearing primary branches (PBPB) (%), Leaf length (LL) (cm), Leaf width (LW) (cm), Leaf area (LA) (cm²), Canopy diameter (CD) (cm), Bean length (BL) (mm), Bean width (BW) (mm), Bean thickness (BT)(mm), Fruit length (FL) (mm), Fruit width (FW) (mm) Fruit thickness (FT) (mm), Clean bean yield (YLD) (Kg/ha)

and Coffee Leaf Rust (CLR).

Analyses of Variance

Analysis of variance (ANOVA) of RCBD was used to see variability using proc mixed procedure of SAS version 9.0 software package [19] (Table 2). Random model which included genotype and location as random factor and genotype \times location interaction was used following statistical model: $Y_{ijk} = \mu + G_i + L_j + B_k (L_j) + GL_{ij} + \epsilon_{ijk}$. Where, Y_{ijk} was the observation for genotype 'i' at location 'j' in replication 'k'. In the model ' μ ' was the overall mean ' G_i ' the effect of the genotype 'i', ' L_j ' was the effect of environment 'j', ' B_k ' block effect, ' GL_{ij} ' the interaction between genotype and location or environment and ' ϵ_{ijk} ' was the random error associated with the k^{th} observation on genotype 'i' in environment. For combined analysis of variance over locations, the homogeneity of error variance was tested using F-max method of Hartley [20]. Traits that showed heterogeneous error variances were square root transformed before combining.

Table 2. ANOVA table skeleton for combined analysis.

SV	DF	MS	EMS
R (location)	l (r-1)	MSr	$\sigma^2_e + gl \sigma^2_r$
Location	l-1	MSl	$\sigma^2_e + r\sigma^2_{gl} + gr\sigma^2_l$
Genotype	g-1	MSg	$\sigma^2_e + r\sigma^2_{gl} + rl\sigma^2_g$
GxE	(g-1) (l-1)	MSgxl	$\sigma^2_e + r\sigma^2_{gl}$
Error	(r-1) (g-1)l	MSerror	σ^2_e

SV-Source of variance, DF- Degree of freedom, MS-Mean square, EMS-Expected mean square GXL-Genotype by Location interaction, l-location, r-replication and g-genotype, Genotypic and Phenotypic variance:- estimated as Johnson et al. [21]:

$\sigma^2_g = \frac{MSg - MSe}{r}$ for individual location, $\sigma^2_p = \sigma^2_g + \sigma^2_e/r$ (Mse/r) $\sigma^2_g = \frac{MSg - MSgl}{rl}$ for over location, $\sigma^2_{gl} = \frac{MSgl - MSe}{r}$, $\sigma^2_p = \sigma^2_g + \frac{\sigma^2_e}{rl} + \frac{\sigma^2_{gl}}{l}$. Where, σ^2_p = phenotypic variance, σ^2_g = genotypic variance, σ^2_{gl} = variance of genotype \times environmental interaction, σ^2_e = environmental variance (Error mean square), MSg = mean square of genotypes, MSe = mean square of error and r = Number of replications

Phenotypic coefficient of variation (PCV) = $\frac{\sqrt{\sigma^2_p}}{\bar{x}} * 100$,

Genotypic coefficient of variation (GCV) = $\frac{\sqrt{\sigma^2_g}}{\bar{x}} * 100$ and

Environmental coefficient of variation (ECV) = $\frac{\sqrt{\sigma^2_e}}{\bar{x}} * 100$, Where \bar{x} = sample mean

PCV and GCV categorized as low (0-10%), moderate (10-20%) and high (>20) [22].

Estimation of heritability in broad sense:- computed as Allard [23] and, Holland et al.[24].

$Hb^2 = \frac{\sigma^2_g}{\sigma^2_p}$ For individual location, $Hb^2 = \frac{\sigma^2_g}{\sigma^2_p}$ Where $\sigma^2_p = \sigma^2_g + \frac{\sigma^2_e}{rl} + \frac{\sigma^2_{gl}}{l}$ for over location. Heritability classified i low (0-20%), moderate (20-50%) and high (>50%) [25].

Estimation of expected genetic advance:- calculated as suggested by Allard [23] as: GA = (K) (σ_p) (Hb^2) Where, GA = expected genetic advance, σ_p = the phenotypic standard deviation, Hb^2 = heritability in broad sense, K = Selection differential (K = 2.06 at 5% selection intensity).

Genetic advance as percent of Mean - GAM = (GA/X) * 100. Where GA and X represent genetic advance and sample mean. It categorized low (0-10%), moderate (10-20%) and high (>20%) [21].

3. Results and Discussions

Analysis of Variance

The results of most traits from analysis of variance of the two individual location indicated that the existence of significance difference between genotypes at probability level of ($p < 0.05$) and ($p < 0.01$) (Table 3). At Mugi, morphological traits like Plant height (PH), total node number of main stem (TNN), inter node length of main stem (IL), number of primary branch (NPB), number of bearing

primary branch (NBPB), percentage of bearing primary branch (PBPB), fruit thickness (FT) and bean characters revealed highly significantly different between genotypes ($p < 0.01$) (Table 3). Whereas like height up to the first primary branch (HFPB), diameter of primary branch (DM), canopy diameter (CD), number of secondary branch (NSB), average length of primary branch (ALPB), number of node per primary branch (NNPB), leaf and fruit characters and yield showed significantly difference among genotypes ($P < 0.05$) at Mugi.

At Haru, traits like PH, HFPB, TNN, DM, CD, NPB, NSB, NBPB, ALPB, NNPB, leaf width (LW), fruit and beans characters revealed highly significantly different among

genotypes ($P < 0.01$) (Table 3). However, IL, PBPB, leaf length (LL), leaf width (LW) and yield (YLD) showed non-significantly different at Haru. Coffee genotypes did not differ significantly in coffee rust disease infection tolerance at Mugi, but significantly different at Haru. There was significant difference among 26 coffee genotypes at both locations in PH, HFPB, TNN, DM, CD, NPB, NSB, ALPB, NNPB, LW and in fruit (fruit length (FL), fruit width (FW), fruit length (FT)) and in bean (bean length (BL), bean width (BW), bean thickness (BT)) traits indicating the existence of genetic variability among the coffee genotypes included in this study.

Table 3. Analysis of variance of quantitative traits.

Plant	Mugi				Haru			
	Blocks	Genotypes	Error	CV (%)	Blocks	Genotypes	Error	CV (%)
Traits	(df= 2)	(df= 25)	(df= 50)		(df= 2)	(df= 25)	(df=50)	
PH	2382.27*	1226.31**	513.87	11.09	3131.11**	546.53**	139.65	7.76
HFPB	0.65ns	30.72*	14.19	13.47	3.25ns	39.50**	5.12	9.57
TNN	8.33ns	19.92**	8.51	10.2	39.27**	10.67**	1.67	5.68
DM	36.07ns	49.62*	26.05	12.57	59.10**	22.85**	7.03	8.27
IL	1.42*	1.29**	0.43	10.2	0.73ns	0.40ns	0.25	8.4
CD	1133.84*	632.46*	319.76	9.78	424.37*	300.85**	108.52	6.54
NPB	11.81ns	49.33**	21.65	11.81	95.55**	39.74**	8.68	9.16
NSB	9.74ns	180.18*	99.88	20.36	78.65ns	177.79**	50.77	20.38
NBPB	18.81ns	16.41**	6.98	12.84	71.12**	13.39**	4.88	16.13
PBPB	222.29*	129.11**	58.34	14.47	162.68**	47.51ns	29.2	12.72
ALPB	264.49*	125.08*	70.38	9.43	131.68**	73.95**	18.9	5.27
NNPB	7.69ns	6.32*	3.45	9.54	1.23ns	4.22**	1.51	5.55
LL	3.64**	0.82*	0.44	4.34	0.58ns	0.61ns	0.46	4.31
LW	0.12ns	0.19*	0.11	5.19	0.43*	0.27**	0.11	5.35
LA	118.72**	44.95*	22.58	7.31	105.29ns	54.80ns	33.35	8.65
FL	1.60*	1.35*	0.63	5.79	1.54*	1.43**	0.4	4.62
FW	0.14ns	0.36*	0.18	3.89	0.54ns	0.63**	0.17	4.21
FT	0.10ns	0.29**	0.13	3.85	0.63*	0.87**	0.13	3.95
BL	2.32**	0.47**	0.05	3.03	1.10**	0.32**	0.06	3.11
BW	0.45**	0.07**	0.03	3.59	0.24**	0.16**	0.02	2.91
BT	0.04ns	0.12**	0.02	6.34	0.01ns	0.05**	0.01	5.1
YLD	121330.89ns	105462.36**	47060.28	44.63	137784.69**	23374.85ns	21118.03	42.01
CLR1	37.66ns	156.88ns	104.65	62.19	145ns	218.40**	30.63	38.63
CLR	(121.91ns)	(212.66ns)	(146.80)	(98.61)	(113.61ns)	(252.86**)	(45.45)	(80.79)

*, **, *** & ns- represent significant at probability level of 0.05, 0.01, 0.001 and non-significant respectively. CLR1- Arcsine transformed coffee leaf rust data

Combined Analysis of Variance

All traits showed homogenous variance except PH, TNN, ALPB and CLR. Traits showed non-homogenous variance transformed by square transformation method before combined analysis (Table 4). Combined analysis of variance revealed highly significant ($p < 0.001$) for HFPB, TNN, NSB and FW (Table 4) among locations. Traits like PH, DM, CD, NPB, PBPB, IL, NNPB and BT showed highly significant difference (0.01) among locations and trait like ALPB showed significant difference (0.05). In all these traits, means were higher at Mugi than at Haru (Table 5). This is due to the ecological nature of Mugi which is known for its high humidity in most seasons and high temperature relative to Haru location which experienced peak humidity in summer season. Mota et al. [26] suggested that lower temperature would trigger declining growth rate of *Coffea arabica* L. Location effect on CLR, yield, leaf, fruit and bean traits was

non-significant except for FW and BT. In line with the present results, Abdulfeta [10] and Masreshaw [27] reported that the existence of variability among Arabica coffee germplasm which were collected from south western Ethiopia using agronomic traits used in this study.

Our results also demonstrated that Haru and Mugi did little differ in bean yield (346 and 486 kg ha⁻¹ at Haru and Mugi respectively) although non-significant. The highest reductions in means at Haru recorded for plant height (25.5%), total number of nodes on the main stem (TNN) (20.6%), number of secondary branches (28.8%) and NBPB (33.4%). Reductions of 10 to 20 % at Haru as compared to that at Mugi were observed in HFPB (15.5%), CD (13.1%), NPB (18.4%), PBPB (19.5%), and FW (10.1%). As shown in Table 5, for the remaining traits (IL, ALPB, NNPB and BT) reductions in means at Haru were less than 10%.

The difference between the genotypes was significant for

only NNPB, FL, FW, FT, and CLR although the difference between genotypes was significant for 18 and 22 of the 23 traits at Haru and at Mugi, respectively. In the combined analysis these differences were masked by the highly significant GxE interaction against which genotype mean squares were tested. GxE interaction was significant for all traits except for NNPB, LL, FW and CLR. These interactions, against which genotypic effects were tested were large in most of the traits and contributed more than 25% to total treatment sum of square (SS) (G + E + GEI) in 16 of the 23 traits (Table 4). The Genotype contributed more than 25% to 18 of the 23 traits. The genotype played minor role (contributed less than 20%) in determining traits such as PH (11.7%), TNN (13.6%), DM (13.0%) and NBPB (17.1%). These traits were determined mainly by the environment (70.5%, 64.0%, 61.2%, and 71.2%, respectively). The genotype played the major role in the determination of traits such as FL (70.1%), FT (69.0%) and CLR (83.2%).

The highly significant GxE interaction showed the non-correspondence between the performances of genotypes at

the two locations, i.e., inconsistency of performance of genotypes over the two locations. For IL (45.6%), LW (48.5%) and leaf area (LA) (43.8%) GxE made large contribution to treatment SS (Table 4). Under such circumstances selection by mean over locations does not identify genotypes that manifest high performance at both locations. Thus, it seems better to divide coffee growing areas into similar ecologies, some similar to Haru and others similar to Mugi and focus on developing coffee varieties with specific adaptation to these ecologies. In line with this, Fikadu *et al.* (2016) [28] reported that similar result of Gx E interaction using 14 characters used in the present study. For FW where GxE contributed only 8.6% to treatment SS and the genotype was the major determinant of the trait (65.9%). The result also indicated that for NNPB, LL and CLR where GxE contributed less to treatment SS and genotype contribution was the dominant (51.8%, 51.1% and 83.2% respectively), selecting elite genotypes based on mean over the two locations may identify common genotypes that are superior at both locations.

Table 4. Over location combined analysis of variance for quantitative traits.

Traits	MSB (df=4)	MSG (df=25)	Gcont. (%)	MSL (df=1)	Econt. (%)	MSG* (df=25)	L GxEcont. (%)	MSE (df=100)	CV (%)
PH	4.15** (4887.75**)	0.95ns (700.46ns)	11.7	147.36** (105798.44**)	70.5	1.48*** (1072.39***)	17.9	0.46 (326.76)	5.12 (10.23)
HFPB	2.11ns	39.95ns	40.2	730.64***	29.4	30.26***	30.4	9.5	11.95
TNN	0.26** (35.84**)	0.11ns (11.55ns)	13.6	13.19*** (1357.89***)	64	0.18*** (19.04***)	22.4	0.05 (5.22)	4.38 (8.91)
DM	0.35* (47.59*)	0.16ns (24.21ns)	13	19.25** (2853.14**)	61.2	0.33*** (48.25***)	25.9	0.11 (18.05)	5.61 (11.69)
IL	2.03*	0.75ns	36.8	9.04**	17.6	0.93***	45.6	0.33	9.36
CD	798.47**	460.09ns	25	22637.13**	49	473.22**	25.7	224.84	17.17
NPB	64.66**	49.39ns	28.9	2045.8**	47.9	39.68***	23.2	15.7	11.08
NSB	41.41ns	187.28ns	28	7770.10***	46.5	170.68**	25.5	74.77	20.58
NBPB	9.00ns	17.68ns	17.1	1841.92**	71.2	12.12**	11.7	7.4	15.88
PBPB	4.05ns	75.90ns	22.2	4150.60**	48.5	100.71**	29.4	50.39	14.9
ALPB	0.59** (198.08**)	0.30ns (105.23ns)	39.8	4.59* (1640.02*)	24.8	0.26** (93.80**)	35.4	0.13 (44.63)	3.92 (7.8)
NNPB	4.46ns	7.21*	51.8	84.10**	24.2	3.33ns	23.9	2.33	7.96
LL	2.81**	0.91ns	51.1	8.76ns	19.7	0.52ns	29.2	0.44	4.41
LW	0.27*	0.24ns	51.2	0.03ns	0.3	0.22**	48.5	0.11	5.24
LA	112**	53.93ns	51.6	120.58ns	4.6	45.83*	43.8	27.94	8.02
FL	1.57*	1.95*	70.1	0.07ns	0.1	0.83*	29.8	0.56	5.5
FW	0.34ns	0.74**	25.6	47.83***	65.9	0.25ns	8.6	0.19	4.13
FT	0.36*	0.83*	69	1.13ns	3.8	0.33***	27.2	0.14	4.02
BL	1.71***	0.46ns	51.8	2.57ns	11.6	0.33***	36.6	0.05	3.11
BW	0.34***	0.14ns	61.5	0.06ns	1.1	0.08***	37.4	0.02	3.26
BT	0.02ns	0.10ns	43	1.67**	27.9	0.07***	29.1	0.02	5.79
YLD	129557.52**	69243.67ns	43.2	766228.09ns	19.2	59593.41*	37.4	34089.15	44.38
CLR	2.33ns (117.75ns)	7.66*** (387.34***)	83.2	0.08ns (1.73ns)	0	1.83ns (78.17ns)	16.8	1.67 (96.13)	53.66 (116.03)

*, **, ***and ns- represent significant different at probability level of 0.05, 0.01, 0.001 and non-significant different respectively. Gcont.- Genotype contribution, Econt. - Environmental contribution and GXEcont.- Genotypes by environmental contribution. PH-Plant height (cm), HFPB-Height up to the first primary branch (cm), TNN-Total node number of main stem, DM-Diameter of main stem (mm), IL-Internodes' length of main stem (cm), CD-Canopy diameter (cm), NPB-Number of primary branch, NSB-Number of Secondary branch, NBPB-Number of bearing primary branch, PBPB-Percent of bearing primary branch, ALPB-Average length of primary branch (cm), NNPB-Number of nodes per primary branch, LL-Leaf length (cm), LW-Leaf width (cm), LA-Leaf area (cm²), FL-Fruit length (mm), FW-fruit width (mm), FT-Fruit thickness (mm), BL-Bean length (mm), BW-Bean width (mm), BT-Bean thickness (mm), YLD-Yield (kg/ha, CLR-Coffee leaf rust (%).

Table 5. Mean separation of quantitative traits among Haru and Mugi locations.

Plant traits												
Locations	PH	HFPB	TNN	DM	IL	CD	NPB	NSB	NBPB	PBPB	ALPB	NNPB
Mugi	204.34a	27.97a	28.61a	40.62a	6.41a	183.47a	39.39a	48.07a	20.57a	52.79a	88.94a	19.49a
Haru	152.26b	23.64b	22.71b	32.06b	5.92b	158.81b	31.83b	32.20b	13.69b	42.60b	82.46b	18.02b
RD at Haru (%)	25.45	15.48	20.62	21.06	7.49	13.13	18.39	28.76	33.41	19.54	7.29	7.54
LSD	7.67	1.30	0.96	1.79	0.24	6.30	1.66	3.63	1.14	2.98	2.98	0.64

Table 5. Continued.

Locations	LL	LW	LA	FL	FW	FT	BL	BW	BT	YLD	CLR
Mugi	15.24a	6.36a	64.99a	13.67a	11.00a	9.25a	7.37a	4.71a	2.23a	486.08a	8.55a
Haru	15.71a	6.33a	66.75a	13.62a	9.90b	9.11a	7.33a	4.76a	2.03b	345.91a	8.34a
RD at Haru (%)	-3.11	0.46	-2.71	0.31	10.06	1.85	-3.49	-0.85	9.31	28.83	2.47
LSD	0.58	0.13	2.22	0.24	0.14	0.16	0.14	0.06	0.04	171.32	4.13

Note-traits assigned by same letter were not significantly different. RD-Reduction, LSD-least significance difference, PH-plant height, HFPB-height up to the first primary branch, TNN-total number of nodes of main stem, DM-main stem diameter, IL-inter node length of main stem, CD-canopy diameter, NPB-number of primary branch, NSB-number of secondary branch, NBPB-number of bearing primary branch, PBPB-percentage of bearing primary branch, ALPB-average length of primary branch, NNPB- number of nodes per primary branch, LL-leaf length, LW-leaf width, LA-leaf area, FL-fruit length, FW-fruit width, FT- fruit thickness, BL-bean length, BW-bean width, BT-bean thickness, YLD-yield and CLR-Coffee leaf rust.

Genotypic and Phenotypic variance at Haru and at Mugi

Results of variability study for 23 traits of coffee at Haru and Mugi were presented in Tables 6 and 7 respectively. At Haru, high phenotypic coefficient of variation (PCV) (>20%) was observed for YLD (25.5%), CLR (110.0%) and NSB (22.0%). These traits had a very wide range; from 172 to 507 kg ha⁻¹ for YLD (a range of 97% of the mean), from 0.4 to 42% infection by CLR (a range of 496% of the mean) and from 26 to 58 secondary branches per tree (a range of 91% of the mean). Moderate PCV (10-20%) was observed for NBPB (15.4%), HFPB (15.4%), and NPB (11.3%); these traits had intermediate range as percent of the mean (60, 63 and 45%, respectively). Our results revealed that for all other 17 traits phenotypic coefficient of variability was low (<10%).

High genotypic coefficient of variation (GCV) (>20%) was observed for only CLR (99.6%). Moderate GCV (10-20%) was observed for NBPB (12.3%), HFPB (14.3%), NPB (10.0%) and for NSB (18.6%). For the remaining 18 traits GCV was low (<10%) indicating that the genotypic variability between the Wollega coffee accessions studied was narrow for most of the traits. This may be due to the fact that these 22 accessions were elite selections from many base collections included in the preliminary screening studies. The present result confirmed with Getachew et al. [34] who reported that low GCV and high PCV for yield traits and low for bean traits.

At Mugi, high PCV (>20%) was observed for bean yield (38.6%) and for CLR (98.4%) likewise at Haru. These two traits had wide range; 257 to 1236 kg ha⁻¹ for bean yield with range of 201.4% of the mean and 1.0 to 37.8% infection by coffee leaf rust with range of 430% of the mean. Moderate PCV (10-20%) was recorded for NBPB, PBPB, PH, HFPB, DM, IL, NPB and NSB whose PCV varied between 10.0% for PH to 15.8% for NSB and whose range as percent of the mean varied from 35.2% for PH to 75.8% for NSB. For the remaining 13 traits PCV was low (<10.0%). High GCV (>20%) was observed for bean yield (28.7%) and for CLR (54.8%). Moderate GCV (10-20%) was recorded only for

NSB (10.2%). Low GCV (<10.0%) was observed for the remaining 20 traits. Thus, similar to the results at Haru there was limited genetic variability for many traits in the Wollega coffee accessions included in this study. In agreement with this, Seyoum et al. [1], Yigzaw [30] and Gizachew et al. [29] found similar result using these quantitative traits (especially for leaf, fruit and bean traits) on *Coffea arabica* L. accessions in Ethiopia.

Broad sense Heritability at Haru and at Mugi

At Haru very high heritability (>80%) was observed in FT, BL, BW, BT, CLR, HFPB and TNN (Table 6). Heritability for these traits was between 80% (BT) to 87% for HFPB. The current result confirmed with the finding of Wagner et al. [31] who reported the similar result using these traits.

High broad sense heritability (Hb) (50-80%) was observed in NBPB, ALPB, NNPB, LW, FL, FW, PH, DM, CD, NPB, NSB and it ranged between 58% for LW to 78% for NPB. Bean yield (10%) and LL (25%) had low and PBPB (39%), LA (39%) and IL (38%) had moderate heritability at Haru. Direct improvement of bean yield is very difficult due to its limited genetic variability and its very low heritability. Thus, indirect selection through traits that are strongly correlated with it and with higher heritability may be a better strategy for improving bean yield at Haru and at locations with similar climatic and edaphic conditions. In line with this, Dawit et al. [32] obtained experimental results that describe the positive correlation between yield and growth traits such as NBPB, ALPB, PH, NNPB, NPB and CD; these indices that the indirect selection of these yield related traits will result coffee yield improvement.

At Mugi heritability was very high (>80%) for BL (89%) and BT (84%) (Table 7). It was intermediate (50 - 80%) for YLD, NBPB, LA, FL, FT, PH, BW, HFPB, TNN, IL and NPB with range of 50% (LA) to 67% (IL). Moderate heritability (20-50%) was observed in ALPB, NNPB, LL, LW, FW, CLR, DM, CD and NSB and it ranged between 31% (CLR) to 49% (CD). Although average heritability was lower at Mugi than at Haru, it was more uniform for the 23

traits; 54+12.5 vs 66.0+21.2, respectively. Bean yield had higher heritability at Mugi than at Haru (55% vs 10%) and the scope of its direct improvement through selection is better at Mugi than at Haru although its indirect improvement through index selection using other correlated traits with higher heritability is better achieved at Haru. This result conformed to the previous result described by Olika *et al.* [11], Lemi and Ashenafi [33] and Getachew *et al.* [34]. Generally, at both locations the low value of Hb indicated that greater value of phenotype variance than genotype variance indicating more influence of environmental factor on those traits. In contrast, high value of Hb showed relatively great influence of genetic factor on those traits indicating the selection of traits for the next breeding program and the possibility of improving genotypes for desired traits [31].

Genetic advance at Haru and at Mugi

Beside the estimated values of GCV and Hb in the

selection process, breeders considered the magnitude of genetic advance (GA) above the population means (GAM) for selection [35]. Hence, from the present study, high GAM (>20%) at Haru was observed in NBPB (20.2%), CLR (101%), HFPB (27.5%) and NSB (32.4%) (Table 6). Moderate GAM (10-20%) was recorded on NNBPB, FT, PH, BT, TNN, DM and NPB which range from 10.3% for NNBPB to 18.2% for NPB. Furthermore, for the remaining 12 traits including bean yield (5.1%) GAM was low (<10%) and ranged between 4.8% for IL to 9.2% for ALFPB. This also shows that direct improvement of bean yield through selection is difficult at Haru. GAM higher than 20% was recorded on bean yield (44.0%) and CLR (62.8%) at Mugi (Table 7). Moderate GAM (10-20%) was recorded for NBPB, PBPB, PH, BT, HFPB, TNN, IL, NPB and NSB with range of 10.6% (TNN) and 15.5% (BT). The remaining 12 traits had low GAM which was between 3.3% for LL and 9.8% for DM.

Table 6. Estimates of variance, coefficients of variations, broad sense heritability and genetic advance of 23 coffee traits at Haru.

CHAR	MIN	MAX	MEAN	RNGE	GENV	EV	PHV	PCV	GCV	H	GA	GAM
YLD	172.36	506.89	345.91	96.71	752.27	7039.34	7791.62	25.52	7.93	0.10	17.56	5.08
NBPB	9.89	18.11	13.70	60.03	2.83	1.63	4.46	15.42	12.29	0.64	2.76	20.19
PBPB	32.59	49.46	42.47	39.72	6.10	9.73	15.84	9.37	5.82	0.39	3.16	7.44
ALPB	71.11	94.67	82.46	28.57	18.34	6.30	24.64	6.02	5.19	0.74	7.61	9.23
NNPB	14.92	19.92	18.02	27.75	1.07	0.33	1.41	6.58	5.75	0.76	1.86	10.34
LL	14.82	16.73	15.71	12.19	0.05	0.15	0.20	2.86	1.42	0.25	0.23	1.45
LW	5.71	7.01	6.33	20.47	0.05	0.04	0.09	4.75	3.61	0.58	0.36	5.65
LA	58.73	77.15	66.75	27.58	7.15	11.12	18.27	6.40	4.01	0.39	3.45	5.16
FL	11.54	15.33	13.62	27.81	0.34	0.13	0.48	5.06	4.30	0.72	1.02	7.52
FW	9.19	11.34	9.90	21.64	0.15	0.06	0.21	4.63	3.94	0.72	0.68	6.92
FT	8.50	10.58	9.09	22.93	0.25	0.04	0.29	5.92	5.46	0.85	0.94	10.37
PH	121.44	170.33	152.26	32.11	135.63	46.55	182.18	8.87	7.65	0.74	20.70	13.60
BL	7.18	8.27	7.63	14.20	0.09	0.02	0.11	4.26	3.86	0.82	0.55	7.22
BW	4.44	5.22	4.76	16.35	0.05	0.01	0.05	4.80	4.50	0.88	0.41	8.69
BT	1.83	2.27	2.03	21.73	0.01	0.00	0.02	6.57	5.88	0.80	0.22	10.83
CLR	3.62	40.26	14.37	254.96	62.59	10.21	72.8	59.38	55.05	0.86	15.12	105.19
HFPB	18.78	33.56	23.64	62.52	11.46	1.71	13.17	15.35	14.32	0.87	6.51	27.53
TNN	17.44	25.44	22.71	35.23	3.00	0.56	3.56	8.31	7.63	0.84	3.28	14.44
DM	25.67	37.31	32.06	36.29	5.27	2.34	7.61	8.61	7.16	0.69	3.93	12.27
IL	5.24	6.86	5.93	27.38	0.05	0.08	0.13	6.16	3.81	0.38	0.29	4.84
CD	137.33	183.67	159.38	29.07	64.11	36.18	100.29	6.28	5.02	0.64	13.19	8.27
NPB	25.22	39.67	32.15	44.93	10.36	2.89	13.25	11.32	10.01	0.78	5.86	18.23
NSB	26.11	57.78	34.97	90.56	42.33	16.93	59.26	22.02	18.61	0.71	11.33	32.40

RNG-Range, GENV-Genotype variance, EV-Environment variance, PHV-Phenotype variance, GCV%-Genotype coefficient of variance, PCV%-Phenotype coefficient of variance, H-Broad sense heritability, GA-Genetic advance, GAM%-Genetic mean advance, PH-Plant height (cm), HFPB-Height up to the first primary branch (cm), TNN-Total node number of main stem, DM-Diameter of main stem (mm), IL-Internodes' length of main stem (cm), CD-Canopy diameter (cm), NPB-Number of primary branch, NSB-Number of Secondary branch, NBPB-Number of bearing primary branch, PBPB-Percent of bearing primary branch, ALPB-Average length of primary branch (cm), NNBPB-Number of nodes per primary branch, LL-Leaf length (cm), LW-Leaf width (cm), LA-Leaf area (cm²), FL-Fruit length (mm), FW-fruit width (mm), FT-Fruit thickness (mm), BL-Bean length (mm), BW-Bean width (mm), BT-Bean thickness (mm), YLD-Yield (kg/ha, and CLR-Coffee leaf rust (%).

Table 7. Estimates of variance, coefficients of variations, broad sense heritability and genetic advance of 23 coffee traits at Mugi.

CHAR	MIN	MAX	MEAN	RNG (%)	GENV	EV
YLD	257.15	1235.94	486.08	201.36	19467.11	15686.75
NBPB	15.56	24.15	20.57	41.80	3.15	2.33
PBPB	41.15	64.10	52.79	43.47	23.59	19.45
ALPB	81.69	111.12	88.94	33.08	18.24	23.46
NNPPB	17.58	23.39	19.49	29.82	0.96	1.15
LL	14.39	16.96	15.24	16.87	0.13	0.15
LW	6.00	7.10	6.36	17.38	0.03	0.04
LA	59.09	73.85	64.99	22.70	7.46	7.53
FL	12.69	15.65	13.67	21.63	0.24	0.21

CHAR	MIN	MAX	MEAN	RNG (%)	GENV	EV
FW	10.41	11.88	11.01	13.36	0.06	0.06
FT	8.03	9.68	9.26	17.76	0.05	0.04
PH	179.89	251.86	204.34	35.22	237.47	171.29
BL	6.70	8.25	7.37	21.10	0.14	0.02
BW	4.43	5.04	4.71	13.04	0.01	0.01
BT	1.93	2.72	2.23	35.05	0.03	0.01
CLR	5.50	36.64	14.21	219.14	17.41	34.88
HFPB	21.33	35.09	27.97	49.20	5.51	4.73
TNN	25.00	37.59	28.61	44.01	3.80	2.84
DM	35.17	52.84	40.62	43.50	7.86	8.68
IL	5.42	7.99	6.41	40.08	0.29	0.14
CD	167.94	220.04	183.47	28.39	104.23	106.59
NPB	33.67	50.82	39.39	43.55	9.23	7.22
NSB	35.00	72.22	49.08	75.84	26.76	33.30

Table 7. Continued.

CHAR	PHV	PCV	GCV	H	GA	GAM
YLD	35153.86	38.57	28.70	0.55	213.89	44.00
NBPB	5.47	11.37	8.63	0.58	2.77	13.48
PBPB	43.04	12.43	9.20	0.55	7.41	14.03
ALPB	41.70	7.26	4.80	0.44	5.82	6.54
NNPB	2.11	7.45	5.02	0.45	1.36	6.97
LL	0.27	3.44	2.36	0.47	0.51	3.33
LW	0.06	3.97	2.60	0.43	0.22	3.52
LA	14.98	5.96	4.20	0.50	3.97	6.11
FL	0.45	4.90	3.58	0.53	0.74	5.39
FW	0.12	3.15	2.20	0.49	0.35	3.17
FT	0.10	3.37	2.53	0.56	0.36	3.92
PH	408.76	9.89	7.54	0.58	24.20	11.84
BL	0.16	5.38	5.09	0.89	0.73	9.91
BW	0.02	3.18	2.40	0.57	0.18	3.74
BT	0.04	8.99	8.22	0.84	0.35	15.48
CLR	52.29	50.89	29.36	0.33	4.92	34.59
HFPB	10.24	11.44	8.40	0.54	3.55	12.69
TNN	6.64	9.01	6.82	0.57	3.04	10.63
DM	16.54	10.01	6.90	0.48	3.98	9.80
IL	0.43	10.22	8.34	0.67	0.90	14.04
CD	210.82	7.91	5.56	0.49	14.79	8.06
NPB	16.44	10.29	7.71	0.56	4.69	11.90
NSB	60.06	15.79	10.54	0.45	7.11	14.49

RNG-Range, GENV-Genotype variance, EV-Environment variance, PHV-Phenotype variance, GCV%-Genotype coefficient of variance, PCV%-Phenotype coefficient of variance, H-Broad sense heritability, GA-Genetic advance, GAM%-Genetic mean advance, PH-Plant height (cm), HFPB-Height up to the first primary branch (cm), TNN-Total node number of main stem, DM-Diameter of main stem (mm), IL-Internodes' length of main stem (cm), CD-Canopy diameter (cm), NPB-Number of primary branch, NSB-Number of Secondary branch, NBPB-Number of bearing primary branch, PBPB-Percent of bearing primary branch, ALPB-Average length of primary branch (cm), NNPB-Number of nodes per primary branch, LL-Leaf length (cm), LW-Leaf width (cm), LA-Leaf area (cm²), FL-Fruit length (mm), FW-fruit width (mm), FT-Fruit thickness (mm), BL-Bean length (mm), BW-Bean width (mm), BT-Bean thickness (mm), YLD-Yield (kg/ha, and CLR-Coffee leaf rust (%))

Genetic advance, Hb and GCV all together provide information of successfully to improve traits of genotypes. When high, Hb, GCV and GAM value combined for desired traits, the involvement of additive gene action expected in that traits. Therefore, at Haru moderate genotypic variance, high heritability and high GAM were observed for NBPB, HFPB and NSB. These traits can be used as indices for improving bean yield which had low heritability and low GAM at Haru. At Mugi bean yield and CLR had high genotypic variance and high genetic advance as percent of the mean although CLR had moderate heritability. Hence, bean yield and CLR can be easily improved via selection at Mugi and at other coffee growing areas with similar soil and climatic conditions. From the population of the top 5% most performing genotypes it is possible to improve yield by 213.89 kg ha⁻¹ at Mugi and CLR

by 5.37% at Mugi and 2.46% at Haru per cycle of selection. Leaf traits (LL, LW, LA), Fruit traits (FL, FW, FT), bean traits (BL, BW), and ALFPB had low GCV and low GAM at both locations. The improvement of these traits seems to be difficult. High Hb with low GAM recorded for many traits expressing traits governed by non-additive gene action. For these traits improvement via cross (hybridization) is better than direct selection. In agreement with the present result, Olika et al. [11], Masreshaw [27] and Gizachew et al. [29] reported similar result for yield and yield related traits used under this study.

4. Conclusion

The current study at both locations indicated the existence of significant difference among tested coffee

genotypes in most traits indicating availability of moderate genetic variability between tested genotypes. The combined analysis of variance of quantitative traits showed significant difference among coffee genotypes only in few numbers of traits although 18 and 22 of the 23 traits showed significant difference among genotypes at Haru and at Mugi, respectively. This was caused by the significant GxE interaction against which genotypes mean square were tested. GxE interaction was significant for all traits except number of nodes per primary branch (NNPB), leaf length (LL), fruit width (FW) and Coffee leaf rust (CLR) indicating non stability performance of coffee genotypes across locations. This indicated that the identification of genotypes with high performance over a wide coffee producing area is very difficult. Hence, it is better to divide coffee growing areas into similar ecologies, some similar to Haru and others similar to Mugi and focus on developing coffee varieties with specific adaptation to these ecologies.

At Haru, moderate genotypic variance, high heritability and high genetic advance as percent of the mean (GAM) were observed for number of bearing primary branch, height up to the first primary branch and number of secondary branch. These traits can be used as indices for improving bean yield which had low heritability and low GAM. At Mugi, bean yield and CLR had high genotypic variance and high GAM; in such condition additive gene action may expect. Hence, Bean yield and CLR can be easily improved via selection at Mugi and at other coffee growing areas with similar soil and climatic conditions.

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